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(54) Title: THROMBUS-SPECIFIC ANTIBODY DERIVATIVES

### (57) Abstract

The invention relates to fibrin-specific single-chain antibodies, thrombolytic agents derived from such antibodies, and DNA fragments coding for such polypeptides. The single-chain antibodies can be used for imaging, while the thrombolytic agents can be used for *in vivo* lysis of thrombi.

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### THROMBUS-SPECIFIC ANTIBODY DERIVATIVES

This invention relates to: novel thrombus (preferably fibrin)-binding molecules ("SCAs"), derived from thrombus (preferably fibrin)-specific antibodies; novel thrombolytic agents ("SCAPAs") derived from SCAs; and genes coding for such SCAs and SCAPAs.

This invention also relates to the uses of the SCAs and SCAPAs and to methods of producing them.

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Background of the Invention

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An injury to a blood vessel normally results in the activation of the complex hemostatic process (Colman et al., 1987) and the formation of a blood clot at the site of injury. The blood clot is predominantly composed of blood platelets emmeshed in a network of fibrin, and its formation protects the organism from bleeding. To restore normal blood flow, the clot is later remodelled and removed by degradation of fibrin by proteolytic enzymes such as plasmin.

Under certain pathological conditions the hemostatic process can also result in the formation of a thrombus, which is a solid mass or plug formed in a living heart or in blood vessels which can cause severe complications (e.g., myocardial infarctions). A thrombosis is due to either local obstruction of blood vessels or distant embolization. The relative amounts of the formative elements of thrombi (i.e., blood platelets, erythrocytes and fibrin) depend on the place in the vascular system where the thrombi are formed (Freiman, 1987).

Fibrin is formed by polymerization of fibrinogen, a protein with a very complex organization. Each fibrinogen molecule is composed of six complexly linked polypeptides (i.e., 2 A-alpha, 2 B-beta and 2 gamma

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polypeptides). The molecule can be divided into a number of domains, the most important of which are the two terminal D-domains and the central E-domain. During fibrin formation, covalent crosslinks are formed between D and D and between D and E domains of adjacent fibrinogen molecules, resulting in interconnected network of fibers with a sufficient mechanical stability to serve as a hemostatic plug (Hantgan et al., 1987). Degradation of fibrin by plasmin results in a variety of degradation products, one of which is the so-called D-dimer which comprises two cross-linked D domains of adjacent fibrinogen molecules.

Therapy of thrombosis and its complications has 15 mainly been directed towards: prevention of local extension of thrombus a by administration anticoagulants such as heparin; and/or acceleration of the rate of dissolution of the thrombus by administration οf thrombolytic agents such as 20 plasminogen activators which are substances initiate formation of plasmin (a proteolytic enzyme that degrades fibrin). The most important plasminogen activators are tissue-type plasminogen activators ("tPA"), urokinase-type plasminogen activators ("uPA") 25 and streptokinase and its derivatives (Bachmann, 1987).

tPA is a protein of 530 amino acids which can be isolated from many tissues but which seems to be mainly produced by endothelial cells as single-chain a polypeptide. The amino acid sequence of this polypeptide and the nucleotide sequence that encodes it have been described by Pennica et al. (1983). tPA is cleaved by plasmin between Arg278 and Ile279 resulting in a molecule consisting of two polypeptide chains held together by one disulfide bridge. In the presence of fibrin both forms are equally active. Structurally, tPA

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can be divided in a finger domain containing lowaffinity fibrin binding sites, a domain with homology
to epidermal growth factor ("EPG"), two kringle domains
with high-affinity fibrin binding sites, and the serine
protease catalytic domain. Because of its binding
capacity to fibrin, plasminogen activation by tPA is
highly fibrin-specific.

An urokinase precursor, designated as prourokinase or single-chain uPA ("scuPA"), is synthesized 10 tissues such as endothelium of the kidney continuous polypeptide of 411 amino acids and a about kiloDalton molecular weight of 54 (kD) acid sequence of ("scuPA-54k"). The amino polypeptide and the nucleotide sequence that encodes it 15 have been described by Holmes et al. (1985). ScuPA is converted to the active form, designated as urokinase or two-chain uPA ("tcuPA") by a proteolytic cleavage plasmin) between Lysine-158 by (e.g., Isoleucine-159 which results in a molecule consisting 20 of two polypeptide chains held together by a single disulfide bond. Three domains can be identified in prourokinase: a EPG domain, one kringle domain and a serine protease catalytic domain. These domains show important homologies with the corresponding domains of 25 tPA. However, neither the EPG domain, nor the kringle domain of scupa have affinity for fibrin.

A low molecular weight form of scuPA ("LMW-scuPA") is formed by an additional proteolytic cleavage between lysine 135 and Lys136, releasing the first 135 amino acids (i.e., the EPG and the kringle domains). The LMW-scuPA is converted to a low molecular weight two-chain urokinase of about 33 kD ("LMW-tcuPA") by a proteolytic cleavage between Lys158 and Ile159.

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European patent publication ("EP") 247674 describes a single-chain uPA of about 32 kD which

apparently is derived from the scuPA by a proteolytic cleavage between Glu143 and Leu144 and which can be recovered in stable conditions and in fair yields from the culture fluid of human lung adenocarcinoma cells of the type CALU-33 (ATCC cell line HTB-55). This form was designated as "scuPA-32k" to differentiate it from the previously known scuPA-54k. ScuPA-32k, like scuPA-54k, can be proteolytically activated, e.g., by plasmin.

Although scuPA-54k does not bind directly to fibrin, it activates fibrin-associated plasminogen much more readily than plasminogen in the plasma (Gurewich et al., 1984). Although scuPA-32k lacks both the EPG and the kringle domain of scuPA-54k, it still displays fibrin selectivity. In contrast, the two active forms, 15 described above, have lost their fibrin selectivity.

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use of all plasminogen activators thrombolytic agents has been hampered by the fact that they will also dissolve existing hemostatic plugs in an organism, thus leading to hemorrhaging. Plasminogen activators that lack fibrin affinity, such will streptokinase and urokinase, also activate circulating plasminogen resulting in impairment of degradation function and platelet of circulating fibrinogen and clotting factors V and VIII (in addition to the fibrin in the thrombus) and leading to systemic effects which have generally been designated as the "lytic state" (Marder and Bell, 1987). In general, it is believed that such systemic effects of plasminogen activator therapy can be minimized by the use of plasminogen activators, such as tPA or scuPA, which demonstrate fibrin-selective plasminogen activation, thus releasing plasmin only in the vicinity of a clot.

tPA and the active forms of uPA have very short circulatory half-lives (5 and 7 minutes for tPA and urokinase, respectively - Sherry, 1987) due to their

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interaction with receptors on cells of the liver. In addition, their activatory half-life is reduced by means of a rapid and irreversible inactivation by plasminogen activator inhibitor I (Haber et al., 1989). Both scuPA-54k and scuPA-32k are resistant to this inhibitor and have longer activatory half-lives. Nevertheless, treatment with any of these agents has had to be over extended periods and in combination with heparin to avoid the occurrence of reocclusion.

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In attempts to improve on the naturally occurring plasminogen activators, several approaches have been followed (see, e.g., Haber et al., 1989; Haber, 1990). Generally, plasminogen activators have been sought which display, in addition to high thrombus (e.g., fibrin)-selective activity, longer circulatory and activatory half-lives.

Antibodies are proteins that are secreted by specialized cells (i.e., the B-lymphocytes) as a part of the immune response of an organism to introduction of foreign molecules (i.e., antigens). Any antibody is highly specific for one particular antigen. With the development of hybridoma technology, monoclonal antibodies, which are highly homogenous with respect to their antigen specificity, could be produced in virtually unlimited quantities (see, e.g., Harlow and Lane, 1988).

The structure of antibodies is well known (see, e.g., Albers et al., 1989). They have a Y-shape and consist of two identical heavy (H) chains of about 440 amino acids and two identical light (L) chains of about 220 amino acids. The various polypeptide chains of a single antibody molecule are connected to each other by four disulfide bridges. In mammals, there are five different classes of antibodies, each of which is characterized by a different type of H-chain (the

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alpha, delta, epsilon, gamma and mu H-chains). addition, antibodies can contain two different L chains (the kappa and lambda L-chains). The type of H-chain effector functions other than defines antigen interactions specificity, such as with antibody receptor molecules on different cells. The difference between the two types of L-chains has not yet been identified. The largest class of circulating antibodies (IgG) have a gamma H-chain.

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10 The Hand L-chains consist of variable and constant domains. The L-chain has a variable domain of about 110 amino acids ("VL") and a constant domain also of about 110 amino acids ("C1"). The H-chain has one variable domain ("V") and three constant domains 15 (" $C_{H1}$ ", " $C_{H2}$ " and " $C_{H3}$ "), each of about 110 amino acids. In each antibody molecule, there are two antiqen binding sites, each of which is formed by the variable domains of one L-chain and one H-chain, so-called complementarity specifically by the 20 determining regions ("CDRs") within the variable domains, and there are three CDRs on each chain. The amino acid sequences of the CDRs are highly variable among antibodies while the sequences of the parts of the variable domains next to and in between the CDRs 25 (i.e., the so-called "framework regions") are much more conserved. The antigen binding site interacts with a well-defined region of the antigen which is designated the "epitope".

The variable domains of the heavy and light chains are encoded by different gene segments which are properly organized in the fully differentiated B-lymphocyte through recombination events. The V<sub>L</sub> region thus consists of a large N-terminal part which is encoded by the so-called variable ("V") gene segment and a short C-terminal part which is encoded by the

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so-called joining ("J") gene segment. The  $V_H$  domain largely consists of a large N-terminal part, which is encoded by a V gene segment, and two smaller parts, encoded by the diversity ("D") gene segment and J gene segment, respectively.

In principle, the  $V_L$  and  $V_H$  are the only components necessary for antigen binding. It has been shown that proteins can be prepared by connecting the nucleotide sequences coding for the  $V_L$  and  $V_H$  regions with a linker sequence coding for a linker polypeptide ("L"), and expression of these hybrid DNA molecules can be obtained in <u>E. coli</u> (Bird et al., 1988; Huston et al., 1988; Chaudhary et al., 1989, 1990). The resulting  $V_L$ -L- $V_H$  or  $V_H$ -L- $V_L$  proteins retained their antigenbinding capacity and can be designated as single-chain antibodies.

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Although general rules have been proposed for designing suitable linkers (see, e.g., PCT publication WO 88/01649), the actual design of a single-chain antibody that retains the affinity and specificity of the original antibody, from which it was derived, and that can be produced in appropriate host cells is far from straightforward.

To increase the fibrin specificity of plasminogen activators, it has been attempted to link the catalytic portions of plasminogen activators with the antigen binding sites of fibrin-specific monoclonal antibodies, for instance by construction of chimaeric molecules with both biological functions. Anti-fibrin antibody 59D8, directed against the amino-terminal six-amino acid sequence of the fibrin beta chain, was chemically conjugated to urokinase and tPA (Bode et al., 1987), and anti-fibrin antibody MA-15C5, directed against human fibrin D-dimer, was conjugated to scupa (Collen et al., 1989; Dewerchin et al., 1990; Collen et al.,

1990). Recombinant DNA technology has also been used to replace parts of the heavy chain of the 59D8 antibody with portions of tPA and uPA catalytic domains (EP 271227, EP 355068 and EP 347078). In general, the results of these attempts have been mixed, and an ideal thrombolytic agent has not yet been identified (Haber et al., 1989; Haber, 1990).

### Summary of the Invention

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invention provides an SCA comprising a This 10 single-chain antibody which can bind in a highly specific manner to at least one thrombus constituent, preferably fibrin. It is preferred that the comprise all or preferably the effective antigenicbinding parts of a monoclonal antibody directed against 15 the thrombus constituent, particularly fibrin, quite particularly fibrin cross-links, in a thrombus. It is particularly preferred that the SCA comprise all or preferably the effective antigenic-binding parts of the light variable domains  $(V_{H})$ and 20 respectively) the monoclonal antibody, linked of through a first linker peptide ("Lab") so as to form a single chain.

This invention also provides a method of using the SCA for imaging of thrombi and for making novel thrombolytic agents comprising the SCA as a thrombus constituent-binding portion, preferably a fibrin-binding portion.

This invention further provides an SCAPA which is a plasminogen activator comprising an SCA as a thrombus-binding portion ("SCA-portion") connected to a plasminogen activating portion ("PA-portion"). It is preferred that the PA-portion comprise at least the catalytic domains of a plasminogen activator, preferably of tPA or uPA, particularly of scuPA. The

C-terminal end of the SCA-portion is preferably directly linked to the N-terminal end of the PA-portion, but both portions can also be linked through a second linker peptide (" $L_{\rm cd}$ ").

5 invention still further provides molecule coding for the SCA ("sca gene") or for the ("scapa gene"), a chimaeric DNA ("chimaeric gene") containing the sca or scapa gene and a vector containing the chimaeric gene. Preferably, the 10 chimaeric gene comprises the following operably linked DNA fragments in the same transcriptional unit: 1) a promoter capable of directing expression of a sca or scapa gene in a procaryotic or eucaryotic host cell; 2) a sca or scapa gene; and 3) suitable 3' regulatory 15 sequences. The chimaeric gene can optionally contain, between DNA fragments 1) and 2), a signal sequence that encodes a polypeptide ("signal peptide") directing the secretion of the SCA or SCAPA from the procaryotic or eucaryotic host.

This invention further provides a method for obtaining the SCA or the SCAPA by: introducing the chimaeric gene in a procaryotic or eucaryotic host cell so that it is actively expressed within the host cell; culturing the host cell; and then recovering the SCA or SCAPA from the culture.

### Brief Description of the Drawings

Fig. 1 - Amino acid sequence of the variable region of the kappa chain of the monoclonal antibody MA-15C5. The numbering of the amino acids follows the generalized numbering described by Kabat et al. (1987). Single lines indicate the borders of the CDR and framework regions.

Fig. 2 - Amino acid sequence of the variable region of the gamma-chain of the monoclonal antibody

MA-15C5. The numbering of the amino acids follows the generalized numbering described by Kabat et al. (1987). Single lines indicate the borders of the CDR and framework regions. The double line indicates the end of the region encoded by the J gene segment.

Fig. 3 - Nucleotide sequence of the cDNA coding for the variable and constant region of the kappa chain of the monoclonal antibody MA-15C5. The amino acid sequence of the variable region is also given (see also Fig. 1). The sequence also comprises the signal sequence. Important restriction sites used during cloning procedures are indicated.

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Fig. 4 - Nucleotide sequence of the cDNA coding for the variable region of the gamma chain of the monoclonal antibody MA-15C5 with the amino acid sequence as given in Fig. 2. The codons for the first four amino acids of the  $V_{\rm L}$  domain are missing. The sequence also comprises part of the coding sequence of the  $C_{\rm H1}$  domain of the MA-15C5 gamma chain. Important restriction sites used during cloning procedures are indicated. The double line indicates the end of the region encoded by the J gene segment.

Fig. 5 - Nucleotide sequence and deduced amino acid sequence of the human uPA cDNA. Important restriction sites used during cloning procedures are indicated. The cleavage site of LMW-tcuPA (single vertical line) and the N-terminus of scuPA-32k (double vertical line) are indicated.

Fig. 6 - Amino acid sequences of preferred SCAs of this invention. The numbers and amino acids in square brackets refer to the amino acid sequence of  $V_L$  of MA-15C5 as given in Fig. 1. The numbers between brackets refer to the amino acid sequence of the first linker peptide  $(L_{ab})$ . The letters and amino acids between accolades refer to the amino acid sequence of

 $V_{\text{H}}$  of MA-15C5 as given in Fig. 2. The sequences derived from the  $V_{\text{L}}$  and  $V_{\text{H}}$  anchor regions are underlined. Residues marked with an asterisk are residues that are mutated with respect to the original sequence.

Fig.7 - Nucleotide sequence of the tac promoter and the PhoA signal sequence. The encoded amino acid sequence of the PhoA signal peptide is also shown. The promoter and signal peptide can for instance be used for the expression and secretion of foreign proteins in <u>E. coli</u>.

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Fig. 8 - A. Nucleotide sequence of the signal sequence of the kappa chain of MA-15C5 monoclonal antibody. The encoded amino acid sequence of the kappa-chain signal peptide is also shown.

- B. Nucleotide sequence of a signal sequence coding for a consensus signal peptide of a human IgG gamma chain. The 11 C-terminal amino acids of this signal peptide are those of the natural signal peptide of the gamma chain of MA-15C5 monoclonal antibody. The signal peptides of Figs. 8a and 8b can, for instance, be used for the expression and secretion of foreign proteins in eucaryotic cells.

Fig. 9 - Nucleotide sequences and corresponding amino acid sequences of the first linker peptide of selected <u>sca</u> genes of the present invention. L<sub>ab</sub>12 (A), L<sub>ab</sub>14 (B) and L<sub>ab</sub>15 (C) respectively correspond to the constructions 1 (and 1A), 5 (and 5A) and 4 (and 4A) (with n=4) of Fig. 6. The sequences of the actual first linker peptides are underlined.

### Detailed Description of the Invention

The single-chain antibody of the SCA of this invention is derived from a monoclonal antibody that is specific for a constituent of thrombi, preferably fibrin, particularly fibrin cross-links. Preferably,

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monoclonal antibody is a murine monoclonal antibody, such as MA-15C5, raised against human fibrin D-dimer. The properties of MA-15C5 have been described by Holvoet et al. (1989), and the construction of recombinant genes coding for the L- and H-chains of MA-15C5, from cDNA libraries of MA-15C5 hybridoma cells, has been described by Vandamme et al. (1990) and in European patent application ("EPA") 90401090.7. Parts of these recombinant genes can be used for the 10 construction of an sca gene. In this regard, the amino acid sequences of the variable domains of the kappa and gamma chains of MA-15C5 are shown in Fig. 1 and Fig. 2, respectively. The CDR and framework regions are these Figures. indicated in The corresponding 15 nucleotide sequences are shown in Fig. 3 and Fig. 4, respectively.

The SCA can have the following general structure  $NH_2-V_1-L_{ab}-V_H-COOH$  or  $NH_2-V_H-L_{ab}-V_L-COOH$ . In order to construct the SCA, the Vu and Vi domains should be 20 linked by an appropriate first linker peptide (Lab). A suitable Lab can be designed using the computerized procedures outlined in PCT patent publication WO 88/01649 (which is incorporated herein by reference). Alternatively, the Lab can be designed by the so-called 25 "spare parts" method as described by Claessens et al. (1989). This method also involves the use, template, of an existing three-dimensional structure of an antibody molecule, with H- and L-chains similar to those of the fibrin-specific monoclonal antibody (e.g., 30 MA-15C5) to be used for the construction of the SCA, to construct a 3D model of the fibrin-specific antibody or at least its framework regions. 3D structures of proteins can be found in, for example, the Brookhaven Database (Bernstein et al., 1977).

The design of the SCA also involves the following three steps:

1) Identifying suitable anchor regions in the  $V_L$  and  $V_R$  domains of the template antibody, to which the first linker peptide  $L_{ab}$  should be attached. The conformation of the anchor regions should be unaffected by introduction of the first linker peptide between them. The attachment sites at the ends of the anchor regions define a gap in which the linker sequence must be placed. The spatial distance between these attachment sites determines a minimum number of amino acids (" $N_{aa}$ ") that are necessary to bridge the gap. Parts of the  $V_L$  and  $V_R$  domains, flanking the anchor regions, can be considered as part of the first linker peptide.

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15 2) Searching a database of 3D protein structures of sufficiently high resolution (e.g., lower than 3 Angstrom) to identify protein fragments, the ends of which overlap the anchor regions and which have the right length and 3D configuration to be able to serve 20 the first linker peptide, bridging the gap identified in step 1. As a general procedure fragments of proteins with a length of Naa to Naa + 6 are assessed as to their suitability as a first linker peptide. The ends of the fragments should fit the anchor regions 25 (e.g., in a least square sense) as closely as possible so that the introduction of the fragment between the anchor regions will not change the correct association of the  $V_i$  and  $V_\mu$  domains. Identification of suitable fragments can be carried out by calculating the root 30 mean square deviations ("rms") between the Cartesian coordinates of the alpha carbon atoms (or the main chain atoms) of the anchor regions and the overlapping regions of the protein fragments (Claessens et al., 1989). Only those fragments, for which the rms falls

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below a certain threshhold determined by the user, are withheld for further study as an  $L_{ab}$ .

- 3) Selecting the most desirable fragments identified in step 2, which preferably conform to the following requirements:
  - The first linker peptide should not interfere with the ordered secondary structure or with the folding of the  $V_L$  and  $V_R$  domains. Secondary structure predictions can be performed according to the procedures described by Jibrat et al. (1987).
  - The regions of the first linker peptide that are exposed to solvent should not contain patches of hydrophobic residues.
- The first linker peptide should be sterically accommodated. Sterical accommodation of fragments can, for instance, be evaluated by calculating the non-bonded energy of the main chain atoms of the linker fragments with respect to the rest of the protein.

The amino acid sequences of the first linker peptide and/or the anchor regions can, if desired, be optimized by introduction of mutations (e.g., substitutions, deletions and/or additions) in order to reduce their non-bonded energy, to minimize their hydrophobicity and/or to improve their flexibility. In this respect, it may be preferred that regions of the first linker peptide, that immediately flank the anchor regions, be mutated to residues that were originally present in the  $V_L$  and  $V_R$  domains.

If no suitable anchor regions can be identified, for instance due to structural constraints or because the gap to be bridged is too big, the  $V_L$  and/or  $V_H$  can be extended by an extension sequence in appropriate

directions, and such a sequence can then serve as an anchor region.

For the construction of an SCA of this invention, it is preferred that the anchor regions be as near as possible to the appropriate C- and N- termini of the  $V_L$  and  $V_H$  or the  $V_H$  and  $V_L$  domains, respectively. However, an anchor region at the C-terminus of the  $V_H$  domain can also be located at the end of the  $V_H$  region that is encoded by the J gene segment.

10 3D structures of proteins (e.g., SCAs) of this be obtained by methods such can invention (Wyckoff al., 1985), crystallography et magnetic resonance spectroscopy (Wüthrich, 1986), structure derivations based on available 3D structures 15 from homologous proteins (see, e.g., Blundell et al., 1987), or from structure predictions based on analysis of the primary structures (for a review, see Taylor, 1988).

The 3D structures of proteins of this invention

20 can be analyzed and modelled by the use of a dedicated computer software package such as the BRUGEL (R) molecular graphics software package (Delhaise et al., 1985 - Plant Genetic Systems N.V., Ghent, Belgium). The effects of substitutions, deletions and additions in known 3D structures or template-derived models on the conformation of the proteins can also be so-analyzed.

In accordance with a preferred embodiment of this invention, the SCA is characterized by an amino acid sequence as shown in Fig. 6. Alternatively, use can be made in such an SCA of a first linker peptide which comes from a naturally-occurring protein and which seems to serve as a natural linker between major functional domains of the naturally-occurring protein, such as a hinge-like sequence of an immunoglobulin.

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For the production of a SCAPA of this invention, the corresponding SCA can be connected to at least the catalytic domain of a plasminogen activator (i.e., the PA-portion). For this purpose, the serine-protease catalytic domain of scuPA (comprising amino acids 144 to 411 in Fig. 5) is preferred, but the catalytic domains of tPA and other plasminogen activators can also be used. The SCA- and PA-portions can, principle, be connected in two ways: the C-terminus of 10 the PA-portion can be linked to N-terminus of the SCA or the C-terminus of the SCA can be linked to the Nterminus of the PA-portion. In order to link the PAportion to the SCA-portion, a suitable second linker peptide (Lrd) should be designed. If the 3D structures 15 of plasminogen activators of interest (or proteins with appreciable homology thereto) are available, this can be done using the same procedure as is used for the construction of the first linker peptide between V, and  $V_{\mu}$  (or vice versa).

If 3D structures are not available for the second linker peptide, use can be made of sequences which seem to serve as natural linkers between major functional domains of the PA-portion. For instance, folding experiments have shown that, in scuPA, the region between Ala132 and Leu144 serves as a natural linker between the kringle and catalytic domains (Oswald et al., 1989). Consequently, when the catalytic domain of scuPA is used for the production of the SCAPA of this invention, in which the C-terminal part of a SCAportion is linked to the N-terminal part of a PAportion, it is preferred that part or all of this region be used as a second linker peptide between the two portions. In such a case, preferred attachment sites on the scuPA are believed to be Ala132, Lys136 and Leu144. In all of these cases, preferred attachment

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sites at the C-terminal part of the  $V_L-L_{ab}-V_H$  of SCAs are believed to be Ser113 (i.e., the end of the  $V_H$  part encoded by the J gene segment), Ser120 (i.e., the actual end of the fourth framework of the  $V_H$  domain), or any other amino acid between these two residues of the heavy chain. A preferred C-terminal attachment site on the  $V_H-L_{ab}-V_L$  of SCAs is believed to be located at Leu104 of the kappa-chain (numbering as in Fig. 1 and Fig. 2).

Alternatively, part of the constant domain following the variable domains of the heavy and light chains of the fibrin-specific antibodies can also be used as the second linker peptide between the SCA- and PA-portions. As another alternative, all or part of the A-domain of scuPA, such as the EGF-like and/or the Kringle domains of scuPA, can be used as the second linker peptide.

It is preferred that proteolytic cleavage of the uPA catalytic domain (e.g., by plasmin or thrombin), resulting in inactive forms of the protein, be prevented. This can be most conveniently done by mutating the amino acids at the cleavage sites so that they are no longer recognized by the proteolytic enzymes. In this regard, Phel57 of scuPA can, for instance, be mutated to Asp157 to remove the Arg156-Phel57 thrombin cleavage site. If also needed, Lys135 of scuPA can be mutated, for instance to Gln135, to remove the Lys135-Gln136 plasmin cleavage site.

The SCA and the SCAPA of this invention can be produced by the expression, in host cells, of the sca and scapa genes, respectively, preferably the chimaeric gene of this invention. The construction of these genes can be achieved in a conventional manner. cDNAs coding for V<sub>1</sub> and V<sub>2</sub> can, for instance, be isolated from cDNA libraries from suitable hybridomas producing thrombus-

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specific, preferably fibrin-specific, antibodies (see, e.g., Vandamme et al, 1990). DNA fragments coding for linkers  $L_{ab}$  and  $L_{cd}$  can be directly synthesized. The gene coding for prourokinase can be obtained as described by Holmes et al. (1985). Appropriate DNA fragments can be ligated to each other by conventional means so as to produce one contiguous DNA fragment coding for the SCA or SCAPA protein of this invention.

The sca and scapa genes can be expressed in 10 suitable procaryotic or eucaryotic host cells placing the genes under the control of a promoter capable of directing their expression in the host cells. Conventional promoters can be used. Preferred promoters for use in E. coli are, for example, 15 regulatable promoters such as: the Ptac promoter (De Boer et al., 1983), the sequence of which is shown in Fig. 7, the P<sub>lac</sub> promoter (Fuller, 1982), the  $P_{trp}$ promoter (Martial et al., 1979), the lambda P, promoter (Bernard et al., 1979) and the  $P_{R}$  promoter (Zabeau and 20 1982). Preferred promoters for mammalian cells have, for example, been described by Menck et al. (1987), Baker et al. (1988), Artelt et al. (1988) and Lee et al. (1981).

If required, a signal sequence can be placed in front of, and in reading phase with, the sca or scapa gene. The signal sequence provides: a) a translation initiation site and b) the necessary functional sequence for exporting the SCA or SCAPA. By signal sequence is meant a DNA fragment coding for fragment ("signal polypeptide peptide") which normally associated with a protein, or subunit of a protein that is translocated out of the cytosol of the host cell -- for example, to the periplasmic space in E. the medium in B.subtilis, coli, to or to endoplasmic reticulum (and, if no other targeting

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information is available, to the medium extracellular space) in eucaryotic cells (or tissues) derived from organisms such as yeasts, insects or mammals. The signal peptide is responsible for the translocation process during which the signal peptide separated or proteolytically removed from the protein or subunit. Signal sequences which can be used are those coding for the signal peptides listed by Watson (1984) or for signal peptides that conform to the general characteristics as outlined by Von Heyne (1988). A preferred signal sequence that can be used in E. coli is the one coding for the phoA signal peptide (Michaelis et al., 1983) which is shown in Fig. 7. Preferred signal sequences, that can be used eucaryotic cells, are those coding for the peptides that are naturally associated with the heavy and light chains of antibodies. In this regard, the amino acid sequence and its encoding nucleotide sequence of the signal peptide of the kappa chain of the MA-15C5 monoclonal antibody is shown in Fig. 8A, and the amino acid sequence and its encoding nucleotide sequence of the signal peptide of the gamma chain of MA-15C5 is shown in Fig. 8B. Another preferred signal sequence is that coding for the signal peptide normally associated with the plasminogen activator, for instance the signal peptide associated with scuPA (Fig. 5 residues Met1 to Gly20 - see also Holmes et al., 1985).

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Although the use of a signal peptide is preferred for the production of the SCA or SCAPA, it is not necessary. A cell can be transformed with just a <u>sca</u> or <u>scapa</u> gene encoding a SCA or SCAPA under the control of a suitable promoter, and the SCA or SCAPA, expressed by the transformed cell intracellularly, can be obtained by lysing the cell.

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Preferred host cells to express the chimaeric gene of this invention are insect cells. In this respect, use can be made of the baculovirus expression vectors (see Maeda, 1989 for a general review). Preferably, the chimaeric gene is placed under the control of the strong polyhedrine promoter, particularly Autographa californica nuclear polyhedrosis virus and expressed in Spodoptera frugiperda using the procedures and vectors described, for example, by Summers and Smith (1987) and Luckow and Summers (1987, 1989) and in 4745051. Other baculovirus US patent expression vectors, such as those described in EP 345152 and EP and PCT publications WO 89/01038 and WO 89/01037, can also be used.

SCA and SCAPA can also be prepared by chimaeric gene capable of being construction of a expressed in other host cells, such as E. coli, B.subtilis, yeasts and mammalian cells (e.g., CHO preferably mammalian cells. Appropriate cells), regulatory promoters, sequences (including regulatory sequences, as well 5' regulatory sequences and enhancer sequences) and, if necessary, signal sequences for such a chimaeric gene are well-known to those skilled in the art.

The chimaeric gene of this invention can be introduced into host cells, the host cells can be cultured, and the SCA or SCAPA can be purified from the host cell culture by conventional means. Secreted SCA or SCAPA can be purified, for example, by affinity chromatography on immobilized epitope (e.g. D-dimer) and/or immunoadsorption to insolubilized antibody raised against the PA-portion.

It goes without saying that the SCA of this invention can be constructed using the variable domain of thrombus-specific, preferably fibrin-specific,

monoclonal antibodies of other than MA-15C5. The corresponding sca gene can be constructed and expressed in analogous ways to those described above. Monoclonal antibodies that can be used are, for instance, those described by Kudryk et al. (1984), Elms et al. (1983), Scheefers-Borchel et al. (1985) and Hui et al. (1986) and in Australian patent publication AU-B-25387/84.

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When the SCA or SCAPA of this invention is used for multiple intravenous applications in patients, it may be preferred to minimize its immunogenicity. This can be achieved (see, e.g., LoBuglio et al., 1989) by replacing the nucleotide sequences coding for the murine framework regions by the corresponding sequences coding for framework regions derived from variable domains of human antibodies as described, for example, by Riechmann et al. (1988) and Verhoeyen et al. (1988) and in EP 328404.

The SCA of this invention can be used for imaging of thrombi. The SCA can be labelled with an opacifying agent, such as an NMR or X-ray contrasting agent, or radioactively labelled in a conventional manner.

The SCAPA of this invention can be used as a thrombolytic agent to treat patients with myocardial infarction, peripheral arterial thrombosis, and stroke, deep venous thrombosis and pulmonary well as embolism. The SCAPA has a number of advantages over The thrombus-specific existing thrombolytics. portion targets the SCAPA, and consequently plasminogen activation activity, to the thrombus. The use of an SCA derived from an antibody specific for fibrin, particularly fibrin cross-links (such MA-15C5), is especially preferred. This ensures that the corresponding SCAPA will remain in contact with the thrombus for a longer period of time during the degradation of fibrin. Because it is believed that the

half-life of the SCAPA is likely to be predominantly determined by its SCA-portion (Collen et al., 1989), it is also expected that the half-life of such a molecule will be greater than its PA-portion alone. believed that the half-life of the SCAPA of this invention also can be increased by producing it: 1) in a non-glycosylated form or in a super-glycosylated form or in a form in which some glycosylation is added to the SCAPA (i.e., to one or more regions of the SCAPA) and other glycosylation is removed from the SCAPA (i.e., from one or more other regions of the SCAPA); and/or 2) in a form which is resistant to plasminogen activator inhibitors; and/or 3) with all or at least a significant part of the A domain (at the N' end of the catalytic domain) of its PA-portion, particularly of scuPA, serving as the second linker peptide. This will permit the application of the SCAPA as a bolus injection and will possibly result in a reduction of the incidence of reocclusion. When compared to other combinations fibrin-specific of antibodies thrombolytic agents, it is also expected that the SCAPA of this invention will display a lower immunogenicity and a better thrombus penetration due to its reduced molecular weight.

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The final conformation of the SCAs and SCAPAs of this invention will depend upon the independent folding of their separate domains  $(V_L, V_H)$ and plasminogen activator catalytic domain) and not upon the association οf disulfide bridges with separate polypeptide chains. It is believed that this will simplify production of these molecules in host cells transformed with sca or scapa genes of this invention as described above. In this regard, it believed that transformed insect or mammalian host cells can properly and secrete the SCAs and SCAPAs of this process

invention, so that they are properly folded for binding to a thrombus constituent and without significant loss of binding activity (as compared to the thrombusspecific antibodies, from which they are derived). 5 Furthermore, the SCAs and SCAPAs of this invention can, if desired, be produced in a glycosylated or superinsect or mammalian cells form in glycosylated transformed with, respectively, sca or scapa genes or mutated sca or scapa genes in which glycosylation sites 10 have been added. Alternatively, the SCAs and SCAPAs can be conveniently expressed in transformed host cells in a non-glycosylated form by mutating the sca and scapa genes at sites which would otherwise encode amino acid sequences which could be glycosylated. In this regard, 15 potential glycosylation sites could be eliminated, for example: in the portion of the nucleotide sequence of Figure 4 encoding the heavy chain domain (VH) of the MA-15C5 antibody, by mutating its AAT nucleotides encoding Asn at positions 261-263 to the nucleotides 20 encoding Asp; and/or in the portion of the nucleotide sequence of Figure 5 encoding scuPA, by mutating its AAT nucleotides at positions 1063-1065 to the nucleotides GAT. In addition, the SCAPAs of this invention can be conveniently expressed in transformed 25 host cells in a form more resistant to a plasminogen activator inhibitor (e.g., PAI-1) by mutating the plasminogen activator catalytic domain encoded by the scapa gene, so that it encodes, for example, a mutant tPA-encoding region as described by Madison et al. 30 1990) or a mutant SCUPA-encoding region in which, from nucleotide 691 to nucleotide 702 in Figure 5, the amino acids Arg Arg His Arg are changed to smaller uncharged amino acids such as Ala or to negatively charged amino acids such as Glu.

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The Examples, which follow, illustrate this invention. Unless otherwise stated in the Examples, all procedures for making and manipulating recombinant DNA were carried out by the standardized procedures 5 described in Sambrook et al, "Molecular Cloning - A Laboratory Manual", Cold Spring Harbor Laboratory (1989). All modelling and analysis of 3D structures of proteins was performed using the BRUGEL(R) software package (Plant Genetic Systems N.V., Ghent, Belgium). 10 All mutagenesis was performed by oligonucleotidedirected construction of mutations by the gapped duplex DNA method (Kramer et al., 1984) using the pMa/c vectors described by Stanssens et al (1987, 1989). Appropriate oligonucleotides were designed according to 15 the general rules outlined by Kramer and Fritz (1988) and synthesized by the phosphoramidite method (Beaucage and Caruthers, 1981) on an Applied Biosystems 380A DNA synthesizer (Applied Bisosystems B.V., Maarssen, Holland). 20

# Example 1 : Design of first linker peptides for the $V_{L}-L_{ab}-V_{H}$ and $V_{H}-L_{ab}-V_{L}$ SCAs

In this and the following examples, the numbering of residues of the MA-15C5  $V_L$  and  $V_R$  domains will follow the standardized numbering of Kabat et al. (1987) (see Figures 1 and 2). The numbering of residues of scuPA will follow that of Holmes et al. (1985) and that of Fig. 5. The numbering of residues of proteins for which the actual 3D structure is known and available in a public database, such as the Brookhaven Database (Bernstein et al., 1977), will follow the numbering as used in this database.

1. Modelling of the 3D structure of the MA-15C5 monoclonal antibody.

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The MA-15C5 antibody contains a kappa light chain and a gamma heavy chain. Thus, the Brookhaven Database was searched for structures of immunoglobulines with similar heavy and light chains. The protein with code pdb2hfl, which is a Fab-lysozyme complex (Sheriff et al., 1987), fulfilled these requirements.

A model of the MA-15C5  $V_L$  and  $V_H$  was obtained by substitution of all residues of the pdb2hfl structure that differed from the MA-15C5  $V_L$  and  $V_H$  sequences with their corresponding residues in the MA-15C  $V_L$  and  $V_H$ .

All substitutions were carried out in the absence of explicit hydrogen (i.e., the no-hydrogen model) and in the absence of water molecules and sequentially from the N- to C-terminus of the  $V_L$  and  $V_H$  domains. Main chain atoms were taken from pdb2hfl template. Side chain orientations were determined by exhaustive map computation varying each of the side chain dihedral angles in steps of 30° in the 0-360 interval and by selecting the configuration with the lowest energy.

The deletions and insertions that would normally be required were not introduced because it was observed that the locations of these mutations were not in regions (i.e., the framework regions) that were important with respect to the linker construction.

## 25 2. Construction of V<sub>I</sub>-L<sub>ab</sub>-V<sub>I</sub>.

While looking for a suitable anchor region at the N-terminus of the MA15C5  $V_{\rm H}$ , it was observed that the first two residues of the gamma-chain had a high temperature factor, thus reflecting mobility. Therefore, these residues were not included in the anchor region. Thus, the anchor region of  $V_{\rm H}$  was defined as the segment comprising residues 3 to 7 (i.e., QLKQS) which forms the end of a  $\beta$ -sheet.

A suitable anchor region at the C-terminus of the MA-15C5  $\,V_L\,$  was found to be the segment comprising

residues 102 to 106 (i.e., TKLEI) which also forms the end of a  $\beta$ -sheet.

The gap between the attachment points of the two linkers is 30.8 Angstrom. Thus, a first linker peptide ( $L_{ab}$ ) of at least 8 amino acids should be sufficient to bridge the gap. Note that there are still two residues (i.e., KR) flanking the C-terminus of the  $V_L$  anchor region, and two residues (i.e., QV) that flank the N-terminus of the  $V_R$  anchor region.

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10 Proteins in the Brookhaven database, that are refined and have a resolution lower than 3 Angström, were searched for fragments which consisted of terminal regions overlapping the anchor regions of  $V_L$  and  $V_R$  and central regions capable of bridging the gap between the 15 attachment points of the anchor regions. The fitting of the fragment terminal regions with the V<sub>1</sub> and V<sub>4</sub> anchor regions was assessed by a least square fit of the atomic coordinates of: 1) the alpha carbon atoms and 2) all main chain atoms (MacLachlan, 1979). This analysis 20 resulted in a root mean square deviation (rms) which should be minimal. The best fragment was found to be the segment comprising residues 22-42 from proteinase K Betzel et al., 1988). (pdb2prk -The following alignment could be made (the anchor regions or the Lab 25 sequence between them are underlined):

... TKLEIKR QVQLKQS... ( $V_L$  gap  $V_H$ )

TYYYDESAGQGSCVYVIDTGI (pdb2prk fragment)

The anchor regions of  $V_L$  and  $V_H$  were, of course, retained. The structure was then subjected to 100 steps of a "Steepest Descent" (Fletcher and Reeves, 1964) energy minimalization procedure fixing all atoms except those of the first linker peptide.

The residues of the first linker peptide that overlap with other regions of  $V_L$  and  $V_H$  were mutated to the residues that were originally present in the  $V_L$  and

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 $V_{\mu}$ . Consequently, the real first linker peptide between the  $V_{t}$  and  $V_{H}$  chains was only 7 residues long (i.e., AGQGSCV). The C residue in this first linker peptide was mutated into a S residue to prevent unwanted disulfide bridge formation. The non-bonded energy of this linker with respect to the rest of the protein was good (E<sub>Van der Wasts</sub>=-39 kcal, E<sub>elec</sub>=-8.5 kcal--both calculated in the no-hydrogen model). The energy of the overall structure was also favorable (E<sub>Van der Waals</sub>=-1355 kcal,  $E_{\text{elec}}$ =-190 kcal--no-hydrogen model). No hydrogen bonds were observed to be formed between the linker and the rest of the structure, and no cavities were Fig. 6 is shown in created. The final SCA (constructions 1 and 1A).

15 The first linker peptide was observed to be located at the side opposite to the antigen-binding site and should not interfere with binding. It was also seen that mutation of the Ile106 and Arg108 residues into Gly or Ser residues also resulted in suitable SCAs (Figure 6, constructions 2, 3, 2A and 3A). Flexibility 20 of the first linker peptide could be increased by replacement of the Q residue in the linker with R followed by 0 to 4 glycine residues (Fig. constructions 4 and 4A) or by replacing the AGQ block 25 of residues in the linker by one or more GGGS blocks of residues.

It was also attempted to use the linker proposed by Bird et al. (1988). This linker has the sequence KESGSVSSEQLAQFRSLD. It was found that the most favorable construction was that in which this linker was attached to Leu104 of  $V_{\rm L}$  and Val2 of  $V_{\rm H}$  (Fig. 6 constructions 5 and 5A).

## 3. Construction of V<sub>H</sub>-L<sub>ab</sub>-V<sub>L</sub>.

As the N-terminus of  $V_{\rm t}$  is located near the CDR region, the first linker peptide should satisfy the

stringent requirement of not interfering with the antigen binding sites. The N-terminus is located at the start of a  $\beta$ -strand situated at the edge of one of two antiparallel  $\beta$ -sheets packed on top of each other. . 5 Interference with the antigen-binding site can be prevented by designing an extension of the V, chain so that the  $\beta$ -sheet at the N-terminus is entered by a  $\beta$ turn preceded by a  $\beta$ -strand. An additional advantage of such an extension is that the gap between the C-10 terminus of the V<sub>H</sub> and the N-terminus of the (extended) V, is smaller so that a shorter first linker peptide immunoglobulins contain a is required. As large collection of turn motifs, the Fab structure (pdb2hfl--Sheriff et al., 1987) was searched for turns 15 for which the main chain heavy atoms of their Cterminal ends could be fitted (in a least square sense) on the main chain heavy atoms of the first three Nterminal amino acids of  $V_i$ . It was observed that the best fit could be obtained by fitting the segment of 20 amino acid residues at positions 68 to 70 of pdb2hFL on the first three amino acids of the V, of MA-15C5 (rms = 0.67 Angstrom). The resulting configuration can be schematically represented as follows (the anchor region and extension are underlined):

25 ....SVTVSS  $\underline{DIK}M...$  (V<sub>H</sub> gap V<sub>L</sub>)
SGSGSGTSY (pdb2hf1 fragment)

Residues 68 to 70 of pdb2hfl are preceded by a  $\beta$ -strand segment (residues 62 to 67). Upon fitting the 68-70 residues of the 62-70 2HFL fragment (i.e. SGSGSGTSY) on the first three residues of the MA-15C5  $V_L$  it was observed that the  $\beta$ -sheet of MA-15C5 was extended by one  $\beta$ -strand.

The actual first linker peptide was then designed between the C-terminus of the  $V_{\rm H}$  domain (using the residues 108-111, i.e., SVTV, as an anchor region) and

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the N-terminus of the extended  $V_L$  domain using the first four residues of the  $V_L$  extension (i.e., SGSG) as an anchor. A search of the 3D structures of proteins for suitable fragments resulted in the identification of a fragment from pdb2sod (superoxide dismutase ("SOD") — Tainer et al., 1982) with 11 residues (SOD residues 038-048 — EGDHGFHVHQF) between the anchor regions. The configuration of the fit can be represented as follows (the anchor regions and first linker peptide are underlined):

... <u>SVTV</u>SS <u>SGSGSGDIKM...</u> ( $V_H$  gap extension- $V_L$ )
TGLTEGDHGFHVHQFGDNT (pdb2sod)

The total first linker peptide (EGDHGFHVHQFSGSGSG) between the original  $V_{\text{H}}$  and  $V_{\text{L}}$  domains is thus composed of this 11 residue pdb2sod fragment plus the six residue pdb2hfl extension sequence which was introduced at the N-terminus of the  $V_{\text{L}}$  domain.

"Steepest Descent" (Fletcher and Reeves, 1964) energy minimalization procedure fixing all atoms except those of the first linker peptide. The first two amino acids of the pdb2sod linker fragment (EG) were then mutated to serine residues to revert to the original V<sub>H</sub> C-terminus. Furthermore, the linker's hydrophobicity was reduced by mutating:

- the F residue at linker position 6 into S
- the V residue at linker position 8 into S
- the F residue at linker position 11 into S

In addition, the S residue at linker position 12 was mutated into a G to increase flexibility. The H residue at linker position 4 was initially not mutated because it was observed that the imidazole was involved in hygrogen bonding with groups in  $V_{\rm H}$ .

The final construction (see also Fig. 6, construction 6) was thus:

{...SVTVSS}-(DHGSHSHQS-GGSGSG)-[DIKM...] corresponding to  $\{V_H\}$ -(linker)- $\{V_L\}$ .

Further modulations of the polarity and hydrophilicity of the first linker peptide can be made by additional modification to it, resulting in the following constructions:

{...SVTVSS}-(DHGSHSEQSGSGSG)-[DIKM...]

{ ... SVTVSS } - (GGGSHSEQSGSGSG) - [DIKM...]

{...SVTVSS}-(GGGSGSGGSGSGSG)-[DIKM...]

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(Fig. 6, constructions 7 to 10). The last of these first linker peptides corresponds to the linker that was used by Huston et al. (1988).

## Example 2 : Construction of sca genes and baculovirus expression vectors containing these genes

The PstI-HindIII fragment of Fig. 4 contains most of the V, domain and part of the N-terminal part of the Cut domain of the MA-15C5 gamma chain. Only the first four amino acids of the V<sub>R</sub> domain are not present 20 (Gln-Val-Gln-Leu). This fragment was cloned into the PstI and HindIII sites of pUC19 (Yanisch-Perron et al., 1985). The Smal-Hind III fragment of the resulting plasmid, pUC19-gamma6 was then cloned in pMc5-8-uts digested with EcoRI, filled in with the Klenow fragment 25 of E. coli DNA polymerase I (Klenow), and further digested with HindIII, yielding plasmid pMc5-gamma6-S. Plasmid pMc5-8-uts can be obtained by cloning a universal translation stop sequence ("uts") with the following sequence : 30

### AGCTTGCTGATTGATTGACCGGATCGATCCGGCT

### ACGACTAACTAACTGGCCTAGCTAGGCCGAGATC

between the HindIII and XbaI sites of the polylinker of pMc5-8 which was described by Stanssens et al. (1987, 1989).

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pMc5-gamma6-S can be used directly for site directed mutagenesis. A stop codon and a EcoRI site was introduced immediately after Ser113 by introduction of the sequence TGAATTC, yielding pMc5-G60-S. The EcoRI sites, and the sequences between them were then deleted by digestion of pMc5-G60-S with EcoRI (filled in with Klenow) and religation. The resulting plasmid was designated as pMc5-G60AE-S.

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chain was obtained The kappa plasmid 10 pcMBDHFR13-15C5KMu (Vandamme et al., 1990). The EcoRI-BglII fragment, shown in Fig. 3, was cloned in the EcoRI and BamHI sites of pMc5-8, yielding plasmid pMc5-Kb. This fragment comprises the signal peptide, the V, domain and the C<sub>1</sub> domain of the Ma-15C5 kappa 15 chain. The EcoRI (filled in with Klenow)-XbaI fragment of pMc5-kb, comprising the kappa chain, was then cloned in the BamHI (filled in with Klenow) and XbaI sites of the baculovirus expression vector pVL1393, yielding pVL1393-K. pVL1393 (now available from British 20 Biotechnology Ltd., Oxford, UK) can be obtained from pVL941, described by Luckow and Summers (1989), by deletion of a 630 bp EcoRI-XmaIII fragment and by extension of the polylinker by insertion of the following sequence in the BamHI site of the pVL941 25 polylinker:

GATCCCGGGTACCTTCTAGAATTCCGGAGCGGCGCCTGCAGATCT
GGCCCATGGAAGATCTTAAGGCCTCGCCGGCGACGTCTAGACTAG

(Summers, personal communication).

The construction of the <a href="mailto:sca">sca</a> gene coding for the <a href="mailto:sca">sca</a> was done as follows. pMc5-G60AE-S was digested with AccI (filled in with Klenow) and XbaI and the fragment, containing the V<sub>H</sub> encoding sequence, was cloned in the StyI (filled in with Klenow) and XbaI sites of pMc5-kb, yielding plasmid pMc5-KG60AE-S. In one mutagenesis experiment, appropriate transition sequences between

the  $V_L$  and  $V_H$  coding regions were then provided. This involved deletion of the remaining part of the kappa constant region, appropriate mutagenesis of the C-terminus of the  $V_L$  and the N-terminus of the  $V_H$  domains (including the addition of the missing N-terminal amino acids of  $V_H$ ), and addition of a linker sequence. Three such mutagenesis experiments, with different linker sequences, led to following three plasmids: pMc5-K12A, pMc5-K14A, pMc5-K15A5 (corresponding to SCAs 1, 5 and 4 respectively in Example 1). These plasmids differ by the first linker peptide between the  $V_L$  and  $V_H$  domains which are shown in Fig. 9.

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The <u>sca</u> genes in pMc5-K12A, pMc5-K14A, pMc5-K15A5 were then introduced in pVL1393-K by replacement of the BamHI-XbaI fragment (coding for the C-terminal end of the MA-15C5 kappa chain) of pVL1393-K with the BamHI-XbaI fragments of pMc5-K12A, pMc5-K14A, pMc5-K15A5, yielding plasmids pVL-K12A, pVL-K14A, pVL-K15A5. These plasmids can be used directly for transfection of insect cells.

# Example 3 : Construction of scapa genes and baculovirus expression vectors containing these genes

The 1475 bp HindIII fragment of the scupa cDNA (Fig. 5) was cloned in the HindIII site of pUC18, yielding plasmid pULscu-PA (Nelles et al., 1987). The NcoI (filled in with Klenow)-HindIII fragment of this plasmid was further subcloned in the BamHI (Klenow) and HindIII sites of pMC5-8, yielding plasmid pMc5-scupa-Nco.

In four subsequent mutagenesis steps carried out on plasmid pMc5-scupa-Nco, the C at position 1356 (Fig. 5) was mutated to a T (resulting in a destruction of a BamHI site), the G at position 966 (Fig. 5) was mutated to an A (resulting in a destruction of a FspI site),

the AAA codon at position 562 (Fig. 5), coding for Lys-135 in scuPA, was mutated to a CAA codon (Gln), and the TTT codon at positions 628-630 (Fig. 5), coding for Phe157 of scuPA, was mutated to a GAT codon (Asp). The resulting plasmid was designated as pMc5-scupa-77-I.

In five additional mutagenesis steps carried out on plasmid pMc5-scupa-77-I, the A at position 648 (Fig. 5) was mutated to a G (resulting in a destruction of an EcoRI site), the G at position 1092 (Fig. 5) was mutated to an A (resulting in the destruction of a PvuII site), the AGGs at positions 691-696 (Fig. 5) were each mutated to a CGT, the G at position 702 (Fig. 5) was mutated to a C (resulting in the creation of a SacII site), and the C at position 624 (Fig. 5) was mutated to a T (resulting in the creation of an StuI site). The resulting plasmid was designated as mPc5-scupa-77-II.

The FspI-XbaI fragments of the pMc5-scupa-77-I and -II plasmids (each comprising the coding sequence of the scuPA catalytic domain) were cloned into the HindIII (filled in with Klenow) and XbaI sites of pMc5-K12A, pMc5-K14A, pMc5-K15A5 (from Example 2), after which the sequences between the C-terminal Ser113 (Fig. 2) codon of the Ma-15C5 V<sub>H</sub> and the N-terminal Ala132 codon of the scuPA were deleted. The resulting plasmids were designated as pMc5-K12A-PA-I, pMc5-K14A-PA-II and pMc5-K15A5-PA-I, pMc5-K12A-PA-II, pMc5-K14A-PA-II

The scapa genes in the pMc5-K12A-PA-I and -II,

pMc5-K14A-PA-I and -II, and pMc5-K15A5-PA-I and -II

plasmids were then introduced in the transfection

vector pVL1393-K (from Example 2) by replacement of a

BamHI-XbaI fragment (coding for the C-terminal part of

the MA-15C5 kappa chain) of pVL1393-K with the BamHI
XbaI fragments of pMc5-K12A-PA-I and -II, pMc5-K14A-

PA-I and -II, and pMc5-K15A5-PA-I and -II, yielding plasmids pVL-K12A-PA-I, pVL-K14A-PA-I, pVL-K15A5-PA-I, pVL-K12A-PA-II, pVL-K14A-PA-II and pVL-K15AS-PAII, respectively. These plasmids can be used directly for transfection of insect cells.

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# Example 4: Expression of sca and scapa genes in insect cells

The sca and scapa genes of Examples 2 and 3 (in plasmids pVL-K12A, pVL-K14A, pVL-K15A5, pVL-K12A-PA-I 10 and -II, pVL-K14A-PA-I and -II and pVL-K15A5-PA-I and introduced and expressed in Spodoptera II) frugiperda (SF9) cells (ATCC no. CRL 1711) using the Autographa californica procedures and polyhedrosis viruses (AcNPV) described by Summers and 15 Smith (1987).

# Example 5 : Purification of SCAs and SCAPAs from insect cell cultures of Example 4

The secreted SCAs expressed in Example 4 are purified by means of affinity chromatography on immobilized fibrin fragment D-dimer.

For purification of secreted SCAPAs expressed in Example 4, this step is followed by immunoadsorption on an insolubilized monoclonal antibody against urokinase, MA-4D1E8, as described by Nelles et al. (1987). The fractions containing urokinase-related antigen are pooled and dialyzed against 0.3 M NaCl, 0.2 M arginine, 0.02 M Tris.HCl buffer pH 7.4, containing 0.01% Tween 80 and 10 KIU/ml aprotinin.

tcuPA is removed from samples equilibrated with dialysis buffer containing 0.2 M arginine by chromatography on benzamidine-sepharose. Fractions devoid of amidolytic activity are pooled.

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Aprotinin is removed by extensive washing on a Centrocon 30 microconcentrator (from Amicon, Danvers, MA, USA).

## 5 Example 6: Purification of SCAPAs from insect cultures of Example 4

The secreted SCAPAs expressed in Example 4 are also purified in a different way from that of Example 5. Each SCAPA is purified by ion exchange chromatography on SP-Sephadex (from LKB, Bromma, Sweden), followed by gel filtration on Sephadex-G100 superfine (from LKB).

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In the first step, 1.5 l of conditioned medium, with a pH adjusted to 5.5, is applied at 4°C and a flow rate of 20 ml/h on a 0.9 x 2 cm SP-Sephadex column equilibrated with 0.05 M NaH2PO4, pH 5.5, containing 0.05 M NaCl, 0.01% Tween 80 and 10 KIU/ml aprotinin. Elution is performed with a 60 ml gradient from 0.05 M to 0.60 M NaCl in 0.05M NaH2PO4, pH 5.5. The fractions containing each SCAPA, as determined with an ELISA specific for uPA-related antigen, are pooled, and the pH is increased to 7.4 with 1 M NaOH. The pooled fractions (representing 7 ml with a concentration of mg of SCAPA per ml) are concentrated on a Centricon 30 microconcentrator (Amicon) to a final volume of 0.5 ml. The concentrated sample is then applied at 4°C and at a flow rate of 4 ml/hr on a 1.0 x 110 cm Sephadex-G100 superfine column equilibrated with 0.02 M Tris-HCl buffer, pH 7.4, containing 0.3 M NaCl, 0.01% Tween 80 and 10 KIU/ml aprotinin. The fractions containing the SCAPA are pooled. Aprotinin is then removed by extensive washing on a Centricon microconcentrator with 0.05 M Tris-HCl buffer, pH 7.4, containing 0.038 M NaCl and 0.01% Tween 80.

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## Example 7: Evaluation of the purified SCAs and SCAPAs from Examples 5 and 6.

Satisfactory antigen-binding activity of the SCAs and SCAPAs of Examples 5 and 6 is found in ELISA using immobilized fibrin fragment D-dimer and rabbit-antimouse antibodies specific for MA-15C5 and goat antibodies specific for total rabbit IgG fraction conjugated to alkaline phosphatase (Voller et al., 1976). Satisfactory urokinase-related antigen activity is also found in ELISA according to Darras et al. (1986).

The SCAs and SCAPAs are also characterized by SDS-PAGE under reducing and non-reducing conditions, and the amino termini of the proteins are determined to verify correct processing. Satisfactory equilibrium association constants of the SCAs for immobilized and dissolved purified fragment D-dimer are determined according to Hogg et al. (1987). SCA is labeled by <sup>125</sup>I to show that there is satisfactory in vitro plasma clot binding capacity (Lijnen et al., 1986) and to determine in vivo half-life.

Satisfactory specific activity of the SCAPAs is shown on fibrin plates (Astrup et al., 1952) comparison with the International Reference Preparation for urokinase (Nelles et al., 1987). The SCAPAs are treated with plasmin (Lijnen et al., 1988) to produce two-chain SCA-tcuPA variants, after which satisfactory amidolytic activity on the synthetic substrate pyroglutamyl-glycyl-arginine-pNA (S-2444 Kabi-Diagnostica) is Satisfactory shown. plasminogen activation activity of the SCAPAs is measured in the presence of an excess of the synthetic substrate S-2251 (Kabi-Diagnostica). Satisfactory in vivo plasma clot binding capacity for 125I-labeled SCAs is shown in a rabbit jugular vein thrombolysis model (Collen et al.,

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1983), and satisfactory in vivo plasma clot lysis activity of the SCAPAs is shown in the quantitative rabbit jugular vein thrombolysis model (Collen et al., 1983) and in the quantitative dog arterial thrombolysis model (Yasuda et al., 1989), in the dog coronary thrombolysis model (Bergman et al., 1983), and in the baboon coronary thrombolysis model (Flameng et al., 1985).

# 10 Example 8: Synthesis and Expression of a pVL-K12A' as in Examples 1 and 2

The computer assisted method of Claessens et al. (1989) was used for the design of a synthetic linker L12 for connecting the carboxyterminal end of the  $\boldsymbol{V}_{L}$ domain of MA-15C5 to the aminoterminal end of its  $V_{\mbox{\tiny H}}$ domain. Since the variable domains of antibodies appear homologous three-dimensional structures, have modeling was based on pdb2hfl (Sheriff et al., 1987). Design of the polypeptide linker was initiated by selecting anchor amino acids (i.e., residues with low reflecting low mobility). factor, temperature suitable anchor region at the carboxyterminus of the  $V_{\rm L}$ domain of MA-15C5 was found to be the segment  $Thr^{102}$ -Lys $^{103}$ -Leu $^{104}$ -Glu $^{105}$ -Ile $^{106}$ , a residues comprising segment that is at the end of a  $\beta$ -sheet. A suitable anchor region at the aminoterminus of the  $V_{\mbox{\scriptsize H}}$  domain of MA-15C5 was found to be the segment comprising residues Gln3-Leu4-Lys5-Gln6-Ser7, a segment that also is at the end of a  $\beta$ -sheet. The attachment sites at the ends of the anchor regions define a gap in which the linker has to be fitted. The 30.8 Å spatial distance between these attachment sites determines a minimum number of amino acids that are required to bridge the gap. This minimum number was found to be 8. The Brookhaven Protein Database was then searched for all peptide sequences

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consisting of 16 to 22 amino acids, so that the length of the peptide linker could be varied from 8 to 14 amino acids. To overlap the  $V_{\mathsf{L}}$  carboxyterminal anchor region, 5 more amino acids were required. To overlap the  $V_{\mbox{\tiny H}}$  aminoterminal anchor region, 3 more amino acids were required. Thus, the number of amino acids had to vary between 16 and 22. This search yielded more than peptide candidates. Secondary structure predictions were then performed according to Jibrat et al. (1987) to select those peptide segments that did not interfere with the ordered secondary structure or with the folding of the  $V_{L}$  or the  $V_{H}$  domain of MA-15C5. In this way, the number of linker peptide candidates was reduced to 82.

15 The fitting of the peptide terminal regions with the  $V_L$  and  $V_{\aleph}$  anchor regions was assessed by a least square fit of atomic coordinates of the  $\alpha$  carbon atoms and of all main chain atoms (MacLachlan, 1979). This analysis resulted in a root mean square deviation that 20 was minimal for a 20 amino acid sequence: Thr-Tyr-Tyr-Tyr-Asp-Glu-Ser-Ala-Gly-Gln-Gly-Ser-Cys-Val-Tyr-Val-Ile-Asp-Thr-Gly-Ile, derived from proteinase K (Betzel et al., 1988). In this sequence, the Thr-Tyr-Tyr-Tyr-Asp fragment overlapped the Thr102-Lys103-Leu104-Glu105-25  ${\rm Ile^{106}}\ {\rm V_L}$  carboxyterminal anchor region, and the  ${\rm Ile-}$ Asp-Thr-Gly fragment overlapped the Gln3-Leu4-Lys5- $GLn^6-Ser^7$   $V_H$  aminoterminal anchor region. The fragments overlapping the anchor regions were mutated to the original anchor region amino acids. The structure was 30 then subjected to 100 steps of a steepest descent energy minimalization procedure (Fletcher and Reeves, 1964), fixing all atoms except those of the linker peptide. From the results of this procedure, it was concluded that the Glu-Ser dipeptide could be replaced by the original  $Lys^{107}-Arg^{108}$   $V_L$  carboxyterminal amino

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acids and that the Tyr-Val dipeptide could be replaced by the original  $Gln^1-Val^2$   $V_H$  aminoterminal amino acids. Thus, a linker peptide with the sequence: Ala-Gly-Gln-Gly-Ser-Cys-Val was derived. In order to prevent unwanted disulfide bridge formation, the peptide linker residue Cys was mutated to Ser, yielding the L12 peptide linker with sequence: Ala-Gly-Gln-Gly-Ser-Ser-Val.

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The cDNA encoding the synthetic peptide linker L12 was then inserted between the cDNA encoding the  $V_{\rm L}$  domain and the cDNA encoding the  $V_{\rm H}$  domain of MA-15C5, resulting in the construction of the synthetic cDNA pMC5-K12A' as described below.

The 419 bp Smal-HindIII fragment from pUC19-G6 15 (Vandamme et al., 1990) was ligated in the EcoRI-HindIII treated pMa/c vector, in which the EcoRI recessing end was filled in with Klenow enzyme, yielding pMA/c-G6. A "TGAATTC" sequence was inserted in pMa/c-G6 by site-directed mutagenesis 20 nucleotides 350 and 351 (on the pUC19-G6 fragment sequence), introducing a TGA STOP codon at the presumed end of the J region of V, and an additional EcoRI site. The resulting plasmid, pMa/c-G60, was digested with EcoRI, treated with Klenow enzyme and religated, 25 yielding pMa/c-Go, in which the EcoRI restriction sites, together with the intervening sequences, are removed.

EcoRI-Bg1II fragment 821 qd The from pCM&DHFR-13-15C5kMu (Vandamme et al, 1990), containing 30 the total kappa chain coding sequence (including the secretion signal) and 3' untranslated sequence, was in EcoRI-BamHI digested pMA/c, inserted pMA/c-Kb. The 406 bp AccI-XbaI restriction fragment from pMA/c-Go, of which the AccI recessing end was made blunt with Klenow enzyme, was transferred to Styl

(filled in)-XbaI treated pMa/c-Kb to yield pMa/c-KGo. In this step, the 226 bp fragment comprising the carboxyterminal part of the kappa constant region (C,) and the kappa 3' untranslated sequence was deleted. A single site-directed mutagenesis with the oligodeoxynucleotide dCAAAGTTGGAAATCAAGCGTGCTGGTCAAGG-CTCTTCTGTTCAAGTTCAGCTGAAGCAGTCAGGACCTGGCC was performed on pMa/c-KG<sub>0</sub> to: i) delete the 328 bp DNA sequence separating the Arg108 of the kappa chain from the codon for Lys5 of the gamma chain; ii) reintroduce cDNA sequence coding for amino acids 1 to 4 missing at the NH2-terminus of V<sub>H</sub>; and iii) insert the peptide linker L12 between the carboxyterminal end of V<sub>L</sub> (Arg<sup>108</sup>) and the aminoterminal end of  $V_H$  (Gln<sup>1</sup>), yielding pMa/c- $K_{12}G_0$ . 3 silent mutations were simultaneously introduced in the kappa coding sequence: the ATA Ile106 codon was changed to ATC; the AAA Lys107 codon was changed to AAG; and the CGG Arg108 codon was changed to CGT. The 818 bp EcoRI (filled in) - XbaI restriction fragment from pMa/c-K<sub>12</sub>G<sub>0</sub> was then transferred to BamHI (filled in)-XbaI treated pVL1393 (British Biotechnology Ltd., Oxford, UK), yielding pVL-K12A'.

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Sf9 cells were grown at 27°C in Grace's insect cell culture medium supplemented with 10% (vol/vol) fetal calf serum, 3.3% (vol/vol) yeastolate, and 3.3% (vol/vol) lactalbumin hydrolysate (TMNF medium) essentially as described by Summers and Smith (1987). The Sf9 cells (2 x  $10^6$  cells in a 25 cm<sup>2</sup> flask) were transfected with 1  $\mu$ g AcNPV DNA and 10  $\mu$ g pVL-K12A' by the Ca-phosphate co-precipitation method (Gorman et al., 1985), and the resulting culture supernatant was harvested 5-7 days later for cloning of recombinant baculovirus and for measurement of human fragment D-dimer binding protein in solid-phase enzyme-linked immunosorbent assay (ELISA).

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For the cloning of recombinant baculovirus, fresh monolayers of Sf9 cells (1.5 x 10<sup>6</sup> Sf9 cells in a 6-well culture plate) were infected with eight 10-fold serial dilutions (between 10<sup>3</sup> and 10<sup>10</sup>) of the cotransfection culture supernatant and subsequently overlaid with 1.5 percent low melting agarose containing 2-fold concentrated Grace's medium. When plaques were well formed (5-10 days post-infection), the putative recombinant plaques (occlusion-negative) were identified using a dissection microscope.

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The recombinant plaques resuspended in 1 ml of TMNF medium and 50  $\mu$ l aliquots were used to infect fresh monolayers of Sf9 cells (2 x 10<sup>6</sup> cells in a 25 cm² culture flask) overlaid with 4 ml TMNF medium. The resulting culture supernatants were harvested 48 h later for assessment of human fibrin fragment D-dimer binding in ELISA. The recombinant virus, AcNpVLK<sub>12</sub>G<sub>0</sub>, was then purified by 4 rounds of plaque purification. For each round, the concentration of fibrin fragment D-dimer binding protein was assessed in ELISA.

The purity of the isolated recombinant virus was confirmed in filter-hybridization experiments (Kafatos et al., 1979). In doing so, the DNA was extracted from 10<sup>6</sup> Sf9 insect cells and x transferred to nitrocellulose. Hybridization was performed with either: a 36 bp probe (GACCCAGTCTCCATCTTCCATGTATGCATCT-CTAGG) complementary to the 72-107 bp aminoterminal sequence of the cDNA encoding the kappa chain of MA-15C5; or a probe complementary to the 134-167 bp polyhedrin cDNA sequence (CTACCCTCGACCCCCAAGACAACTACCTA-GTGGC) that is deleted in the pVL-K12A' cDNA by recombination.

For the large scale production of the SCA encoded by pMC5-K12A' and pVL-K12A', called "scFV- $K_{12}G_0$ ", 40 x  $10^6$  Sf9 cells in 175 cm<sup>2</sup> culture flasks were infected

with 200 x 10<sup>6</sup> plaque forming units of recombinant virus  $AcNpVLK_{12}G_0$ . After incubation for 48 h at 27°C, the conditioned medium, containing up to 15  $\mu g$  scFv- $K_{12}G_0$  per ml, but on average approximately 4.5  $\mu g/ml$ , was removed and centrifuged at 1,000xg for removal of cell debris.

scFv-K<sub>12</sub>G<sub>0</sub> was purified by ion exchange chromatography on SP-Sephadex followed by gel filtration on Sephadex-G100 superfine. In the first 10 step, 1.5 l of conditioned medium with a pH adjusted to 4.5 was applied on a 0.9 x 2 cm SP-Sephadex column equilibrated with 0.05 M NaH2PO4, pH 4.5, containing 0.05 M NaCl, 0.01% Tween 80 and 10 KIU/ml aprotinin. Elution was performed with a 60 ml gradient from 0.05 M 15 to 1.0 M NaCl in 0.05 M NaH2PO4, pH 5.5. The fractions containing scFv-K<sub>12</sub>G<sub>0</sub>, as measured in ELISA specific for fibrin fragment D-dimer binding protein, were pooled and the pH was increased to 7.4 with 1M NaOH. The pooled fractions (representing 7 ml with 20 concentration of 0.7 mg scFv-K<sub>12</sub>G<sub>0</sub> ml) per were concentrated on Centricon 10 microconcentrator а (Amicon) to a final volume of 0.05 ml. The concentrated sample was applied on a 1.0 x 110 cm Sephadex-G100 superfine column equilibrated with 0.02 M Tris-HCl 25 buffer, pH 7.4, containing 0.3 M NaCl, 0.01% Tween 80 10 KIU/ml aprotinin. The fractions containing scFv-K<sub>12</sub>G<sub>0</sub> were pooled and found to migrate to a single 25,500 Mr band on reduced SDS gel electrophoresis.

 $m NH_2$ -terminal amino acid analysis of the scFv- $K_{12}G_0$ , so obtained, revealed that the MA-15C5 kappa signal peptide was cleaved off by the insect cells just in front of mature kappa Asp¹ residue. The scFv- $K_{12}G_0$  was also found to bind to immobilized D-dimer with an affinity constant of 4 x  $10^9$  M¹, as compared to 2.0 x  $10^{10}$  m¹ for intact MA-15C5. This finding indicates

that, in scFv- $K_{12}G_0$ , the MA-15C5  $V_L$  and  $V_H$  domains can reassociate efficiently, resulting in the reconstitution of an intact, functionally active, antigen binding site. Further, it can be concluded that, provided the first linker peptide which connects the  $V_L$  and  $V_H$  domains does not put any spatial distance or structural constraints on the overall  $\beta$ -sheet structure of the framework regions, the molecular interactions responsible for the conserved framework structure will assure the proper folding of the hypervariable domain loops in the antigen binding site in the Fv fragment.

When injected as a bolus (2.8  $\mu$ g/kg), scFv-K<sub>12</sub>G<sub>0</sub> was cleared from the plasma of rabbits with a half-life of 10 minutes and a clearance rate of 5.1 ml/min<sup>-1</sup>, as compared to 90 minutes and 210 ml/min<sup>-1</sup> for intact MA-15C5. These results indicate that scFv-K<sub>12</sub>G<sub>0</sub> can be useful for targeting radioisotopes or plasminogen activators to blood clots in vivo.

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## Example 9: Synthesis and Expression of a pVL-K12A-PA-II' as in Example 3

A transfer vector pVL-K12A-PA-II', encoding the SCAPA called " $K_{12}G_0S_{32}$ ", for expression in Sf9 insect cells was constructed starting from the plasmids pMA/c- $K_{12}G_0$  and pVL-K12A' of Example 8, pULscu-PA (Nelles et al., 1987) and the pMA/c mutagenesis vector. A 1117 bp NcoI-HindIII fragment from pULscu-PA, containing the sequence encoding a scuPA fragment consisting of amino acids 67 to 411, connected to the 3'-untranslated sequence, was ligated in the BamHI-HindIII treated pMa/c mutagenesis vector, yielding pMa/c-scu-PA'.

By site-directed mutagenesis, the following mutations were introduced in the DNA sequence of scuPA

(nucleotide numbering refers to Fig. 5; amino acid numbering refers to the Holmes et al. (1985) scuPA The Lys<sup>135</sup>-Lys<sup>136</sup> sequence) to yield pMa/C-scu-PA'm. plasmid cleavage site in urokinase was removed by 5 Lys<sup>135</sup> substituting with Gln, using the 17-mer oligonucleotide dGGGCTTTTGTCCATCTG (underlined residue differs from the wild type residue). The Arg156-Phe157 thrombin cleavage site in urokinase was removed by mutating Phe<sup>157</sup> to Asp (nucleotides 628-630) with the 10 31-mer oligonucleotide dCCCAATAATCTTATCGCGAGGCCTCAGAG-TC. To facilitate the screening of the mutants, a StuI restriction site (nucleotides 619-624) simultaneously created by changing the CCC Pro155 codon to CCT. The 33-mer oligonucleotide 15 dGACAGAGCCCCCGCGGTGACGACGGTAGATGGC was used to modify 3 codons: Arg178 and Arq<sup>179</sup> AGG rare (nucleotides 691-696) were replaced by CGT codons, while for screening purposes, the Arg181 CGG codon was changed to CGC, generating a SacII restriction site 20 (nucleotides 699-704). The BamHI restriction site in urokinase (nucleotides 1352-1357) was deleted changing the ATC Ile399 codon to ATT with the 18-mer oligonucleotide dGTGACTGCGAATCCAGGG. This mutation was performed to facilitate further manipulation of the 25 chimeric cDNA, using the BamHI restriction site present in the variable kappa light-chain coding sequence. One of the two FspI restriction sites (nucleotides 963-968) was removed by changing the GCG Ala369 codon to GCA with the 20-mer oligonucleotide dCGGGATGGCTGTGCACACCT. The 30 FspI enzyme cleaves the remaining site precisely in front of amino acid Ala 132, which was used as the NH2amino acid of the truncated terminal facilitate further manipulation of the chimeric gene, one of the two EcoRI restriction sites (nucleotides 646-651) was deleted by changing the GAA Glu163 codon to

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GAG with the 30-mer oligonucleotide dGATGGTGGTGAACTCTCCCCCAATAATCTT, and the PvuII restriction site (nucleotides 1090-1095) was removed by changing the CAG Gln<sup>311</sup> codon to CAA with the 19-mer oligonucleotide dGTCATTTTCAGTTGCTCCG.

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The 613 bp BamHI-HindIII fragment from pMa/c-K<sub>12</sub>G<sub>0</sub>, which encodes the carboxyterminal sequence of scFv-K<sub>12</sub>G<sub>0</sub> and of which the HindIII end was filled in with Klenow enzyme, was ligated in BamHI-FspI treated pMa/c-scu-PA'm. The resulting plasmid pMa/c-12VS contained the sequence encoding the carboxyterminal region of scFv-K<sub>12</sub>G<sub>0</sub> in front, but out of frame, of the aminoterminal sequence of the truncated catalytic domain of scuPA. oligonucleotide-directed mutagenesis Deletion performed on pMa/c-12VS to delete the 22 nucleotides that still separated the carboxyterminal amino acid (Ser<sup>232</sup>) of scFv- $K_{12}G_0$  and the first amino acid (Ala<sup>132</sup>) of the truncated catalytic domain of scuPA, yielding  $pMa/c-12G_0S_{32}$ . The 51-mer oligonucleotide **dagaggaggcttttgtccatctgctgaggagacggtgactgaggttccttg** used was complementary to the 9 carboxyterminal amino and to the 8 acids of the scFv-K<sub>12</sub>G<sub>0</sub> molecule aminoterminal amino acids of the low molecular weight form (truncated catalytic domain) of scuPA. Finally, to reconstruct the total scFv-K12G0, domain of the chimeric the BamHI-XbaI fragment of molecule, containing the carboxyterminal K<sub>12</sub>G<sub>0</sub> coding sequence, was replaced by the 1521 bp BamHI-XbaI restriction fragment from pMA/c-12G<sub>0</sub>S<sub>32</sub>, yielding pVL-K12A-PA-II'.

Sf9 cells were grown at 27°C in Grace's insect cell culture medium supplemented with 10% (vol/vol) fetal calf serum, 3.3% (vol/vol) yeastolate, and 3.3% (vol/vol) lactalbumin hydrolysate (TMNF medium), essentially as described by Summers and Smith (1987). The Sf9 cells (2  $\times$  10 $^6$  cells in a 25 cm $^2$  flask) were

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transfected with 1  $\mu$ g AcNPV DNA and 10  $\mu$ g pVL-K12A-PA-II' by the Ca-phosphate co-precipitation method (Gorman et al, 1985), and the resulting culture supernatant was harvested 5-7 days later for cloning of recombinant baculovirus and for assessment of human fibrin fragment D-dimer binding protein in solid-phase enzyme-linked immunoassay (ELISA).

For the cloning of recombinant baculovirus, fresh monolayers of Sf9 cells (1.5 x 106 Sf9 cells in a 6-10 well culture plate) were infected with eight 10-fold 10<sup>3</sup> and dilutions (between 10<sup>10</sup>) cotransfection culture supernatant and subsequently overlaid with 1.5% low melting agarose containing 2fold concentrated Grace's medium. When plaques were 15 well formed (5-10 days post-infection), the putative recombinant plaques (occlusive-negative) identified using a dissection microscope (Summers and Smith, 1987).

The recombinant plaques were resuspended in 1 ml of TMNF medium, and 50 ul aliquots were used to infect fresh monolayers of Sf9 cells (2 x 10<sup>6</sup> cells in a 25 cm<sup>2</sup> culture flask) overlaid with 4 ml TMNF medium. The resulting culture supernatants were harvested 48 h later for assessment of human fibrin fragment D-dimer binding protein in ELISA.

The recombinant virus (AcNpVLK<sub>12</sub>G<sub>0</sub>S<sub>32</sub>) was then purified by 4 rounds of plaque purification. For each round, the expression of fragment D-dimer binding protein and of uPA-related antigen was assessed in ELISA. The purity of the isolated recombinant virus was confirmed in filter-hybridization experiments (Kafatos et al., 1979).

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For the large scale production of  $K_{12}G_0S_{32}$ , 40 x 10<sup>6</sup> Sf9 cells in 175 cm<sup>2</sup> culture flasks were infected with 200 x 10<sup>6</sup> plaque forming units of recombinant virus

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AcNpVLK<sub>12</sub>G<sub>0</sub>S<sub>32</sub>. After incubation for 48 h at 27°C, the conditioned medium was removed and centrifuged at 1,000 g for removal of cell debris.

 $K_{12}G_0S_{32}$  was purified as described in Example 6 by ion exchange chromatography on SP-Sephadex followed by gel filtration on Sephadex-G100 superfine.

The specific activity of the resulting  $K_{12}G_0S_{32}$  towards a chromogenic substrate for urokinase was  $\leq$  1,000 IU/mg before and 100,000 IU/mg uPA equivalent after conversion to its two-chain derivative with plasmin. The specific activity of both the single-chain and two-chain form on fibrin plates was 100,000 IU/mg uPA equivalent. Activation of plaminogen by  $K_{12}G_0S_{32}$  obeyed Michaelis-Menten kinetics with Km=  $2.9 \pm 0.6 \mu$ M and a  $k_2$ =  $3.7 \pm 0.6 s^{-1}$  (mean  $\pm$  SD; n= 3), as compared to Km=  $12 \mu$ M and  $k_2$ =  $4.8 s^{-1}$  for recombinant scuPA-32k (low  $\underline{M}_r$  scuPA consisting of amino acids Leu<sup>144</sup> to Leu<sup>411</sup>).

Single-chain  $K_{12}G_0S_{32}$  induced a dose- and timedependent lysis of a <sup>125</sup>I-fibrin labeled human plasma clot immersed in citrated human plasma; fifty percent lysis in 2 h was obtained with 0.70  $\pm$  0.07  $\mu$ g/ml (mean  $\pm$  SD; n= 5) as compared to 8.8  $\pm$  0.1  $\mu$ g/ml for recombinant scuPA-32k (mean  $\pm$  SD; n= 3).

With two-chain  $K_{12}G_0S_{32}$ , fifty percent clot lysis in 2 h required 0.25  $\pm$  0.03  $\mu$ g/ml (mean  $\pm$  SD; n= 3) as compared to only 0.62  $\pm$  0.04  $\mu$ g/ml (mean  $\pm$  SD; n= 2) for recombinant tcuPA-32k. Fragment D-dimer inhibited the fibrinolytic activity of  $K_{12}G_0S_{32}$  (50 percent inhibition with 6  $\mu$ g fragment D-dimer/ml) but not of scuPA-32k.

These results indicate that low  $\underline{M}_r$  scuPA of this invention can be targeted to a fibrin clot with a single-chain Fv fragment of a fibrin-specific antibody, resulting in a 13-fold increase of the fibrinolytic

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potency of the single-chain form and a 2.5-fold increase of the potency of the two-chain form, as compared to that of their uPA-32k counterparts.

Needless to say, this invention is not limited to the transformation of a specific host microorganism or for this purpose, of a chimaeric the use, containing any specific promoter, signal sequence, sca scapa gene and/or 3' transcription regulation sequence of this invention, or the use of any specific SCA or SCAPA, expressed by such a transformed host, for the specific purposes mentioned above. In this regard, equivalents of the foregoing Examples will be readily apparent to those skilled in the art in view of the disclosure herein of the invention. For example, the DNA sequences of the described sca and scapa genes (and consequently the amino acid sequences of the resulting SCAs and SCAPAs) can be easily modified by: replacing some codons with others that code either for the same amino acids or for other amino acids; and/or 2) deleting or adding some codons; provided that such modifications do not substantially alter the biological properties of the encoded SCAs or SCAPAs.

Also this invention is not limited to an SCA or an SCAPA derived from a monoclonal antibody directed to fibrin or fibrin D-links, such as MA-15C5 antibody. This invention encompasses SCAs and SCAPAS derived from antibodies directed to other thrombus monoclonal constituents such as: a) antibodies to blood platelets, for example antibodies to resting and activated e.g., receptors, antibodies surface platelet platelet membrane glycoprotein IIb/IIIa (Bode et al., 1990) or antibodies (e.g., MA-libs-1) specific for ligand-occupied receptor conformers (Frelinger et al., 1990); or b) antibodies to alpha 2-antiplasmin. This invention also encompasses SCAs and SCAPAs derived from WO 91/16353 PCT/EP91/00767

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other monoclonal antibodies directed to fibrin such as the 59D8 antibodies (Bode et al., 1987).

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#### CLAIMS

- A single-chain antibody: which is specific to a constituent, preferably thrombus fibrin; preferably comprises all or especially a part of a monoclonal antibody directed against the thrombus constituent, especially fibrin, particularly fibrin cross-links, quite particularly fibrin D-dimer; which is preferably properly folded for binding to the constituent; and/or which thrombus is preferably glycosylated; the part of the monoclonal preferably comprising all or especially a part of variable domains of the light and heavy chains of the monoclonal antibody, connected by means of a first linker peptide.
- 2. The single chain antibody of claim 1 which is derived from monoclonal antibody MA-15C5 and which preferably has one of the amino acid sequences shown in Fig. 6.
- 3. A thrombolytic agent, comprising the single-chain antibody of claim 1 or 2, connected, preferably by means of a second linker peptide, to a plasminogen activating portion, preferably a catalytic domain of urokinase or tissue plasminogen activator, preferably of scuPA.
- 4. The thrombolytic agent of claim 3, which has increased half-life and which: a) is in a non-glycosylated form or in a super-glycosylated form or in a form in which some glycosylation is added to, and other glycosylation is removed from, the thrombolytic agent; and/or b) is in a form which is resistant to a plasminogen activator inhibitor; and/or c) has all or at least a significant part of the A domain of its

plasminogen activating portion serving as the second linker peptide.

- 5. The thrombolytic agent of claim 3 or 4 in which: a) the plasminogen activating portion is the catalytic domain of urokinase, preferably having the amino acid sequence shown in Fig. 5 from Leu144 to Leu411; and b) in which the C-terminal end of the single-chain antibody is either directly connected to the N-terminal end of the catalytic domain or preferably is connected by the second linker peptide, derived from the urokinase region connecting the kringle and catalytic domains, preferably having the amino acid sequence shown in Fig. 5 from Ala132 to Glu143.
- 6. A DNA fragment coding for the single-chain antibody of claim 1 or 2 or the thrombolytic agent of anyone of claims 3-5.
- A chimaeric gene for transforming a host cell to express the DNA fragment of claim 6, comprising the following operably linked DNA fragments in the same transcriptional unit: i) promoter capable а directing the expression of the DNA fragment in the host cell, preferably an insect cell, a mammalian cell, or Escherichia coli cell, particularly an insect cell or a mammalian cell; ii) the DNA fragment of claim 6; iii) a suitable 3' transcription regulation sequence for the host cell; and optionally, between fragments i) and ii), iv) a signal sequence coding for a signal peptide capable directing secretion of the expression product of the DNA fragment ii) from the host cell.
- 8. A host cell, preferably an insect cell, a mammalian cell, or Escherichia coli cell, particularly

an insect or mammalian cell, transformed with the chimaeric gene of claim 7.

- 9. A method for producing the single-chain antibody of claim 1 or 2 or the thrombolytic agent of anyone of claims 3-5, comprising: culturing the host cells of claim 8; and recovering the antibody or thrombolytic agent from the culture medium.
- 10. A process, comprising the use of the single-chain antibody of claim 1 or 2 for imaging of a thrombus in a living organism.
- 11. A process, comprising the use of the thrombolytic agent of anyone of claims 3-5 for dissolving a thrombus in a living organism.
- 12. A vector for transforming the cell of claim 8, comprising the chimaeric gene of claim 7.

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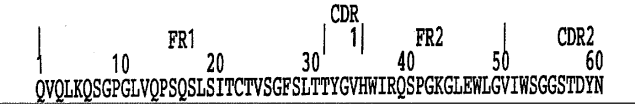
FIGURE 1

	10	FR1	20	ı	CDR1 30		FR 40	. <del>-</del>	5	CDR2		FR:
DIKMTQSE	PSSM:	YASLGE	RVTVT	CKA	SQDIN	SYLSW:	IQQKP	GKSPK	(TLIY	RGNRL	VAC	VPS
		F	R3		İ	CDR3		FR4	i			
	70	-	80		90	02110	100		110	)		
RFSGSGSG	QDY.	SLTISS	LEYED	VGV	YYCLRY	DEFPI	TFGS	GTKLE	IKR			

No insertions

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#### FIGURE 2



CDR2 FR3 CDR3 FR4 70 80 90 100 110 120 AAFISRLSINKDNSKSQVFFKMQANDTAIYYCARNYWGTSDYWGQGTSVTVSSAKTTPPS

### <u>Insertions</u>

in FR3: 82ABC = 82NSL

in CDR3: 100ABCDEFGHIJK = 100-----M

EgoDT	10			20		3	0		40			50
ECORI GAATTCO	GCA	CATG	AGGA	CC C	CTGC	TCAG	T TT	CTTG	GAAT	CTI	GTTG	CTC
TGGTTTC		GTAT(			GAC	ATC	AAG	ATG	ACC	CAG		
98 CCA TCT Pro Ser		ATG	TAT	GCA	TCT	CTA	GGA	GAG	AGA	GTC		
134 GTC ACT Val Thr	TGC	AAG	GCG	AGT	CAG	GAC	ATT	AAT	AGC	TAT	TTA	
ם	179 amHI			188			197			206		
AGC TGG Ser Trp	ATC	CAG Gln	CAG Gln	AAA Lys	CCA Pro	GGG Gly	AAA Lys	TCT Ser	CCT Pro	AAG Lys	ACC Thr	
215 CTG ATC Leu Ile	TAC Tyr	CGT	GGA	AAC	AGA	TTG	GTT	GCT	GGG	GTC Val	CCA Pro	
251 TCA AGG Ser Arg	TTC Phe	260 AGT Ser	GGC	AGT	269 GGA Gly	TCT	GGG	278 CAA Gln	GAT	TAT Tyr	287 TCT Ser	

## FIGURE 3 (continued 1)

· ·		296	1.00	100	305	63.0	mam	314	a m	OTHOR	323	<b>⊘</b> mm	
CTC	ACC	ATC	AGC Ser	AGC	Leu	GAG	TAT	GAA Glu	Asp	Val	GGA Gly	Val	
TAT Tyr	TAT	TGT	CTA Leu	CGG	TAT	GAT	GAG	TTT	CCA	TTC	ACG Thr	TTC Phe	
GGC	TCG	GGG	377 ACA Thr	AAG	TTG	GAA	ATA	AAA	CGG	GCT	ATG(	l05 TG	
CAC	CAACI	115 IGT 1	ATCC!	42 ATCTT	5 C C(	CACCA	435 ATCCA	GTO	GAGCA	445 GTT	AACA	45 TCTGG	5
GGT(	GCT(	165 CAG !	rcgt(	47 STGCT	5 T C1	TGAA	485 CAAC	TTC	TACC	495 CCA	AAGA	50 CATCA	5 A
TGT(	E CAAGI	515 EGG <i>1</i>	AAGA1	52 TGAT	5 G <b>G</b> (	AGTO	535 AACG	ACA	AAAT	545 GGC	GTCC	55! TGAAC	5
GTT(	5 GACT	65 GA 1	rcago	57 Jacag	5 C AA	AGAC	585 AGCA	CCT	'ACAG	595 CAT	GAGC	605 AGCACO	) ~ ~
CTCA												655 CTGTGA	
GGC(	6 CACTO	65 AC <i>P</i>	AAGAC	67 ATCA	5 A CT	TCAC	685 CCAT	TGT	CAAG	695 AGC	TTCA	705 ACAGGA	)

### FIGURE 3 (continued 2)

715	725	735	745	<b>75</b> 5
ATGAGTGTTA	GAGACAAAGG	TCGGGCGAGC	TCGAATTAAT	TCACTCCTCA
765	775	785	795	805
GGTGCAGGCT	GCCTATCAGA	AGGTGGTGGC	TGGTGTGGCC	AATGCCCTGG
815	825			
	Bqll	<u>[]</u>		
CTCACAAATA	CCACTGAGAT	CT		

D est	LŦ	10	,		2	0		2	9		3	8
CTG	CAGG	AAT (	g aa	G CA	G TC	A GG	A CC	T GG	C CI	A GT	G CA l Gl	G
CCC Pro	TCA Ser	CAG	AGC	CTG	TCC	ATC	ACC	TGC	ACA	GTC	74 TCT Ser	GGT
TTC Phe	83 TCA Ser	TTA Leu	ACT Thr	92 ACC Thr	TAT Tyr	GGT Gly	101 GTA Val	CAC His	TGG Trp	110 ATT Ile	CGC Arg	CAG Gln
TCT	CCA	GGA	AAG	GGT	CTG	GAG	TGG	CTG	GGA	GTG	ATA Ile	TGG
AGT Ser	GGT Gly	GGA	AGC Ser	ACA	GAC	TAT	AAT	GCA	GCT	TTC	191 ATA Ile	TCC Ser
AGA Arg	CTG	AGC	ATC Ile	AAC	AAG	GAC	AAT	TCC	AAG	AGC	CAA Gln	GTT Val
TTC	TTT Phe	AAA	ATG	AAC	AGT	CTG	CAA	GCT	AAT	GAC	ACA Thr	GCC

## FIGURE 4 (continued 1)

ATA TAT Ile Tyr	281 TAC TGT Tyr Cys	290 GCC AGA Ala Arg	AAT	TAT	299 TGG Trp	GGA	ACC Thr	308 TCT Ser	ATG MET
317 GAC TAC Asp Tyr	TGG GGT Trp Gly	326 CAA GGA Gln Gly	ACC Thr	335 TCA Ser	GTC Val	ACC Thr	344 GTC Val	TCC Ser	TCA Ser
353	362		371		38	30	r.	39( :oRI	)
GCC AAA Ala Lys	ACG ACA Thr Thr	CCC CCA Pro Pro	TCT Ser	GTCI	TATCO	CA CI	GGAA	TTC	j

400 <u>HindII</u>] ATATCAAGCTT

	20	30	40	50
HindIII AAGCTTCGGG	CCAGGGTCCA C	CTGTCCCCG CA	GCGCCGTC GCG	CCCTCCT
GCCGCAGGCC	70 ACCGAGGCCG C	80 CGCCGTCTA GC	90 GCCCCGAC CTC	99 GCCACC
ATG AGA GCC	117 CTG CTG GCG Leu Leu Ala	CGC CTG CTT	CTC TGC GTC	CTG
GTC GTG AGC	153 GAC TCC AAA Asp Ser Lys	GGC AGC AAT	GAA CTT CAT	CAA Gln
GTT CCA TCG	189 AAC TGT GAC Asn Cys Asp	TGT CTA AAT	GGA GGA ACA	TGT
GTG TCC AAC	AAG TAC TTC Lys Tyr Phe	TCC AAC ATT	CAC TGG TGC	AAC
261 TGC CCA AAG Cys Pro Lys	270 AAA TTC GGA Lys Phe Gly	279 GGG CAG CAC Gly Gln His	288 TGT GAA ATA Cys Glu Ile	GAT Asp
AAG TCA AAA	306 ACC TGC TAT Thr Cys Tyr	GAG GGG AAT	GGT CAC TTT	333 TAC Tyr

## FIGURE 5 (continued 1)

342  CGA GGA AAG GCC AGC ACT GAC ACC ATG GGC CGG Arg Gly Lys Ala Ser Thr Asp Thr MET Gly Arg  378  CTG CCC TGG AAC TCT GCC ACT GTC CTT CAG CAA Leu Pro Trp Asn Ser Ala Thr Val Leu Gln Gln	Pro Cys  ACG TAC  Thr Tyr
CTG CCC TGG AAC TCT GCC ACT GTC CTT CAG CAA	ACG TAC Thr Tyr
TIEM TIO TIP USH DOT UTW THE AUT DOG OTH OTH	450
414 423 432 441 CAT GCC CAC AGA TCT GAT GCT CTT CAG CTG GGC His Ala His Arg Ser Asp Ala Leu Gln Leu Gly	CTG GGG
AAA CAT AAT TAC TGC AGG AAC CCA GAC AAC CGG Lys His Asn Tyr Cys Arg Asn Pro Asp Asn Arg	AGG CGA
CCC TGG TGC TAT GTG CAG GTG GGC CTA AAG CCG Pro Trp Cys Tyr Val Gln Val Gly Leu Lys Pro	CTT GTC
531 540 549 558 FSpI	567
CAA GAG TGC ATG GTG CAT GAC TGC GCA GAT GGA Gln Glu Cys MET Val His Asp Cys Ala Asp Gly	AAA AAG
CCC TCC TCT CCT CCA GAA GAA TTA AAA TTT CAG Pro Ser Ser Pro Pro Glu Glu Leu Lys Phe Gln	603 TGT GGC

## FIGURE 5 (continued 2)

	612			621			630			639		
CAA	AAG	ACT	CTG	AGG	CCC	CGC Arg	TTT	AAG	ATT	ATT	GGG	GGA
GAA	TTC	ACC	ACC	ATC	GAG	666 AAC Asn	CAG	CCC	TGG	TTT	GCG	GCC
ATC	TAC	AGG	AGG	CAC	CGG	GGG Gly	GGC	TCT	GTC	ACC	TAC	GTG
TGT Cys	GGA	GGC	AGC	CTC	ATC	AGC Ser	CCT	TGC	TGG	GTG	ATC Ile	AGC Ser
GCC	ACA	CAC	TGC	TTC	ATT	783 GAT Asp	TAC	CCA	AAG	AAG	GAG	GAC
TAC Tyr	ATC Ile	GTC	TAC	CTG	GGT	CGC Arg	TCA	AGG	CTT	AAC	TCC	AAC
ACG	CAA	GGG	GAG	ATG	AAG	TTT Phe	GAG	GTG	GAA	AAC	CTC Leu	ATC Ile

## FIGURE 5 (continued 3)

	882	ብአብ -	አአፖ	891	መአሶ	አርሮ	900 CCT	ርአ <u></u> ሮ	እሮር	909 Ста	GCT	<sub>ር</sub> ያር	918 Cac
	Leu	His	Lys	Asp	Tyr	Ser	Ala	Asp	Thr	Leu	Ala	His	His
			927		#	936			945			954	
	AAT Asn	GAC Asp	ATT Ile	GCC Ala	TTG Leu	CTG Leu	AAG Lys	ATC Ile	CGT Arg	TCC Ser	AAG Lys	GAG Glu	GGC Gly
		-			972								
		1	'SDL		-							3 m/1	maa
	AGG	TGT	GCG Ala	CAG	Pro	Ser	Ara	ACT	ATA Ile	Gln	ACC Thr	Ile	Cvs
	-	_					_						
	999			800			017	666	<b>.</b>	1026	000	101	1035
	CTG	CCC	TCG	ATG MRT	TAT	AAC Agn	GAT Acn	Pro	CAG	TTT Phe	GGC Gly	ACA Thr	AGC
	пси				_		-						
			044		1	053	<b></b>	1	1062		1	071	<b></b>
	TGT	GAG	ATC	ACT	GGC	TTT	GGA	AAA	GAG	AAT	TCT Ser	ACC Thr	GAC
	Cys	GTU	116	TIII	GTÅ	rne	GIY	пÃр	GTU	וומת	DCT	TIIT	nah
		1080			089		1	098		1	107		
	TAT	CTC	TAT	CCG	GAG	CAG	CTG	AAA	ATG	ACT	GTT	GTG	AAG
	Tyr	Leu	Tyr	Pro	Glu	Gin	Leu	Lys	MET	'I'nr	Val	vai	гĀг
1	116		1	125		1	134		1	143		1	152
	CTG	ATT	TCC	CAC	CGG	GAG	TGT	CAG	CAG	CCC	CAC	TAC	TAC
	Leu	Ile	Ser	His	Arg	Glu	Cys	GIN	Gin	PTO	His	Tyr	Tyr

## FIGURE 5 (continued 4)

	•	1161		•	1170			1179			1188		
GGC Glv	TCT	GAA	GTC	ACC	ACC	AAA	ATG	CTG Leu	TGT	GCT	GCT	GAC	
•	1197			1206			1215			1224		•	
CCA	CAG	TGG	AAA	ACA	GAT	TCC	TGC	CAG Gln	GGA	GAC	TCA Ser	GGG Gly	
1233	000	ATTA	242	mem	maa	1251	<b>733</b>	aaa	260	1MC	3 (7III)	1269	
								GGC Gly					
ልርጥ	GGA	1278 ልጥጥ	GTG	AGC	287 TGG	GGC	CGT	1296 GGA	TGT	GCC	305 CTG	AAG	
Thr	Gly	Ile	Val	Ser	Trp	Gly	Arg	Gly	Cys	Ala	Leu	Lys	-
•	1314		1	323		1	332		1	341			
								GTC Val					
1350		1 BamHI			1	368		1	377		1	386	
CCC	TGG	ATC	CGC	AGT	CAC	ACC	AAG	GAA	GAG	TAA	GGC	CTG	
PIO	Trp	116	Arg	261	nis	TIII	тÃ2	Glu	GIU	ASII	GTÄ	теп	
_	CTC	TGA						5 YG GG					
WTQ	Leu	•											

## FIGURE 5 (continued 5)

1445	1455	1465	1475	1485
		GAGTCATCTC		
1495	1505	1515	1525	1535
CAGAGACACT	AACGACTTCA	GGGCAGGGCT	CTGATATTCC	ATGAATGTAT
1545	1555	1565	1575	1585
CAGGAAATAT	ATATGTGTGT	GTATGTTTGC	ACACTTGTTG	TGTGGGCTGT
1595	1605	1615	1625	1635
GAGTGTAAGT	GTGAGTAAGA	GCTGGTGTCT	GATTGTTAAG	TCTAAATATT
1645	1655	1665	1675	1685
TCCTTAAACT	GTGTGGACTG	TGATGCCACA	CAGAGTGGTC	TTTCTGGAGA
1695	1705	1715	1725	1735
GGTTATAGGT	CACTCCTGGG	GCCTCTTGGG	TCCCCCACGT	GACAGTGCCT
1745	1755	1765	1775	1785
GGGAATGTAC	TTATTCTGCA	GCATGACCTG	TGACCAGCAC	TGTCTCAGTT
1795	1805	1815	1825	1835
TCACTTTCAC	ATAGATGTCC	CTTTCTTGGC	CAGTTATCCC	TTCCTTTTAG
		1865 CTGGGTGGGG		

### FIGURE 5 (continued 6)

1895 1905 1915 1925 1935 GAATATTTAT ATTTCACTAT TTTTATTTAT ATTTTGTAA TTTTAAATAA

1945 1955 1964 AAGTGATCAA TAAAATGTGA TTTTTCTGA

V<sub>L</sub>-L<sub>ab</sub>-V<sub>H</sub> constructions

Construction 1.
[D1...G99-SGTKL-EIKR]-(AGQGSSV)-{QV-QLKQS-G8...S113}

Construction 2. [D1...G99-SGTKL-EG\*KG\*]-(AGQGSSV)-{QV-QLKQS-G8...S113}

Construction 3.
[D1...G99-SGTKL-ES\*KS\*]-(AGQGSSV)-{QV-QLKQS-G8...S113}

Construction 4.
[D1...G99-SGTKL-ES\*KS\*]-(AGR(nG)GSSV)-{QV-QLKQS-G8...S113} with n=0 to 4

Construction 5
[D1...K103-L]-(KESGSVSSEQLAQFRSLD)-{V-Q3...S113}

Construction 1A.
[D1...G99-SGTKL-EIKR]-(AGQGSSV)-{QV-OLKQS-G8...S120}

Construction 2A. [D1...G99-SGTKL-EG\*KG\*]-(AGQGSSV)-{QV-QLKQS-G8...S120}

Construction 3A.
[D1...G99-SGTKL-ES\*KS\*]-(AGQGSSV)-{QV-QLKQS-G8...S120}

Construction 4A.
[D1...G99-SGTKL-ES\*KS\*]-(AGR(nG)GSSV)-{QV-QLKQS-G8...S120} with n=0 to 4

Construction 5A
[D1...K103-L]-(KESGSVSSEQLAQFRSLD)-{V-Q3...S120}

#### FIGURE 6 (continued 1)

### V<sub>H</sub>-L<sub>ab</sub>-V<sub>L</sub> constructions

Construction 6. {Q1...T107-SVTV-SS}-(DHGSHSHQSGGSGSG)-[DIK-M4...R108]

Construction 7. {Q1...T107-SVTV-SS}-(DHGSHSEQSGGSGSG)-[DIK-M4...R108]

Construction 8. {Q1...T107-SVTV-SS}-(GGGSHSEQSGGSGSG)-[DIK-M4...R108]

Construction 9.
{Q1...T107-SVTV-SS}-(GGGSGSGGSGGSGSG)-[DIK-M4...R108]

**EcoRI** 

GAATTCGAGCTCGAGCTTACTCCCCATCCCCTGTTGACAATTAATCATCGG

CTCGTATAATGTGTGGAATTGTGAGCGGATAACAATTTCACACAGGAAACA

BamHI BamHI

GGATCCGCGGATCCGTGGAGAAAATAAA GTG AAA CAA AGC ACT ATT Met Lys Gln Ser Thr Ile

GCA CTG GCA CTC TTA CCG TTA CTG TTT ACC CCT GTG ACA Ala Leu Ala Leu Leu Pro Leu Leu Phe Thr Pro Val Thr

AAA GCG Lys Ala

A.

ATG AGG ACC CCT GCT CAG TTT CTT GGA ATC TTG Met Arg Thr Pro Ala Gln Phe Leu Gly Ile Leu

TTG CTC TGG TTT CCA GGT ATC AAA TGT Leu Leu Trp Phe Pro Gly Ile Lys Cys

B.

ATG GCT GTC TTA GGG CTG CTC TTC TGC CTA GTG Met Ala Val Leu Gly Leu Leu Phe Cys Leu Val

ACA TTC CCA AGC TGT GTC CTA TCC Thr Phe Pro Ser Cys Val Leu Ser

A. Lab 12

CA AAG TTG GAA ATA AAA CGG GCA GGC CAA GGG AGC TCA
Lys Leu Glu Ile Lys Arg Ala Gly Gln Gly Ser Ser

47 56 65

GTA CAA GTA CAA CTA AAG CAG TCA GGA CCT GGC C
Val Gln Val Gln Leu Lys Gln Ser Gly Pro Gly

B. Lab 14

CG GGG ACA AAG TTG AAA GAA TCA GGA TCA GTC TCG AGT Gly Thr Lys Leu Lys Glu Ser Gly Ser Val Ser Ser

GAA CAA TTA GCA CAA TTT AGA TCT TTA GAT GTA CAG CTG Glu Gln Leu Ala Gln Phe Arg Ser Leu Asp Val Gln Leu

AAG CAG TCA GGA CCT GG Lys Gln Ser Gly Pro

#### FIGURE 9 (continued 1)

C. L<sub>ab</sub>15

CG GGG ACA AAG TTG GAA TCA AAG AGC GCT GGC CGC GGC Gly Thr Lys Leu Glu Ser Lys Ser Ala Gly Arg Gly

GGC GGC GGC TCG AGT GTC CAA GTA CAG CTG AAG CAG Gly Gly Gly Ser Ser Val Gln Val Gln Leu Lys Gln

83 TCA GGA CCT GGC Ser Gly Pro Gly

#### INTERNATIONAL SEARCH REPORT

International Application No PCT/EP 91/00767

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) <sup>6</sup>							
According to International Patent Classification (IPC) or to both National Classification and IPC							
IPC <sup>5</sup> :	C 07 K 15/28, A 61 K 37/	02, C 12 N 15/13, C	12 P 21/08				
II. FIELDS	S SEARCHED	-					
Minimum Documentation Searched 7							
Classification	on System	Classification Symbols					
IPC <sup>5</sup>	C 07 K, C 12 N, C	12 P, A 61 K					
	Documentation Searched othe to the Extent that such Documen	r than Minimum Documentation its are included in the Fields Searched <sup>a</sup>					
III. DOCU	MENTS CONSIDERED TO BE RELEVANT		1				
Category *	Citation of Document, 11 with indication, where a	ppropriate, of the relevant passages 12	Relevant to Claim No. 13				
x	EP, A, 0271227 (THE GEN 15 June 1988	NERAL HOSPITAL CORP.	1,3,6-9,12				
	Soo Pullegraps	, <u>.</u> .	! 2				
Y			<b>2</b>				
Y	of monoclonal antib	"Binding properties codies against human f cross-linked fibrin	2				
A	model in rabbits", see the whole artic  EP, A, 0347078 (CELLTEG 20 December 1989 see page 4, lines 2	pages 307-313 cle CH LTD)	1-9,12				
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"A" doc con "E" earir filin "L" doc white cita "O" doc othe "P" doc late	ument defining the general state of the art which is not sidered to be of particular relevance iier document but published on or after the international g date ument which may throw doubts on priority claim(s) or ch is cited to establish the publication date of another tion or other special reason (as specified) ument referring to an oral disclosure, use, exhibition or armeans ument published prior to the international filing date but r than the priority date claimed	"X" document of particular relevant cannot be considered novel or involve an inventive step "Y" document of particular relevant cannot be considered to involve document is combined with the ements, such combination being of	e; the claimed invention out or theory underlying the e; the claimed invention cannot be considered to e; the claimed invention in inventive step when the or more other such docubivious to a person skilled				
	FICATION  Actual Completion of the International Search	Date of Mailing of this International Sec	erch Report				
	th August 1991	1 1 SEP 1991'					
Internation	sternational Searching Authority Signature of Authorized Officer						
EUROPEAN PATENT OFFICE			AZELAAS				

Form PCT/ISA/210 (second sheet) (January 1985)

FURTHER	INFORMATION CONTINUED FROM THE SECOND SHEET					
A	Trends in Biotechnology, vol. 6, no. 2, February 1988, Elsevier Publications, (Cambridge, GB),	1-9,12				
	G. Williams: "Novel antibody reagents: production and potential", pages 36-39,42	·				
	see page 39: "Single chain antibodies"	,				
		, .				
F	·					
<b>v</b> . □ ов	SERVATION WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 1					
This Internati	onal search report has not been established in respect of certain claims under Article 17(2)(a) for the follow	wing reasons:				
	numbers 10, 11 because they relate to subject matter not requirity, namely:	ired to be searched by this				
t .	PCT-Rule 39.1 (iv): methods for treatment of the human	·				
or	animal body by surgery or therapy as well as diagnostic					
met	hods					
2. Claim	numbers because they relate to party of the leterantions	d annii andura dhat da aat aa aa				
	numbers  because they relate to parts of the international he prescribed requirements to such an extent that no meaningful international search can be carried out, s	pecifically:				
	numbers because they are dependent claims and are no	t drafted in accordance with				
the second and third sentences of PCT Rule 6.4(a).						
VI.□ OB	SERVATIONS WHERE UNITY OF INVENTION IS LACKING 2					
This Internati	onal Searching Authority found multiple Inventions in this International application as follows:					
		ļ				
1 As al	required additional search fees were timely paid by the applicant, this international search report covers a	all searchable claims				
of the International application						
2 As or	ly some of the required additional search fees were timely paid by the applicant, this international search i claims of the international application for which fees were paid, specifically claims:	report covers only				
	tains					
3 No re	quired additional search fees were timely paid by the applicant. Consequently, this international search rep					
then	vention first mentioned in the claims, it is covered by claim numbers:	on is restricted to				
4 As al	searchable claims could be searched without effort justifying an additional fee, the International Searching	Authority did not				
Remark of	payment or any additional fee					
	iditional careb face were accommon to the second					
. —	ditional search fees were accompanied by applicant's profest plest accompanied the payment of additional search fees					
	prystown or maintains destruit ites					

#### ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

EP 9100767 SA 46812

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 06/09/91

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

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